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Evaluation of Calibration Parameters and Performance of the Video

Imaging Technique of Assessing Exposure (VITAE System)

by

Keith M. Groth

A thesis submitted in partial fulfillment of the requirements for the degree of

Master of Science

University of Washington

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CHAPTER I

INTRODUCTION

1.1 Need for the Capability to Assess Occupational Dermal Exposure

Assessing exposures to chemicals that can be absorbed through the skin presents the industrial hygienist with a perplexing and sometimes frustrating dilemma. The fact that three of the articles appearing in the February 1993 AIHA Journal addressed surface wipe sampling is a testament to an increasing concern over the impact of dermally absorbed chemicals. In addition to increased awareness of percutaneous absorption as a potentially major factor in total body burden, this concern probably is fueled by frequent reductions in the allowable levels of many airborne contaminants for the workplace. As allowable workplace levels drop, the percentage of total body burden that is a result of dermal absorption may increase. This is especially true if dermal exposures are not controlled as rigorously as inhalation exposures. In fact, in many instances where the chemical has a low vapor pressure and is not aerosolized, dermal absorption is the primary route of exposure (McArthur, 1992).

Although percutaneous absorption of workplace chemicals has been recognized as a very real problem, methods for evaluating these exposures are still in their infancy. In a recent OSHA rule governing the use of a suspected carcinogen that is dermally absorbed (4,4'-methylene-dianiline), OSHA conceded that in certain situations 95% of exposure comes from dermal absorption. However, OSHA failed to establish a dermal exposure limit, citing difficulties in quantifying exposures, correlating amount absorbed with risk, and inability to select a reliable biological indicator as reasons why such a limit is currently infeasible (OSHA,

1992). If an organization with the resources of OSHA, is unable to determine an appropriate method of monitoring dermal exposures to a chemical such as MDA, an independent industrial hygienists attempting to do so is faced with a daunting task.

1.2 <u>Current Methods for Assessing Dermal Exposures</u>

Currently, the sampling methods that are available to assess dermal exposures include surrogate skin, surface contamination, skin contamination, and biological. Each of these methods have inherent limitations. Used in combinations of two or more, they may be employed to sufficiently assess dermal exposures. But in some instances, as noted in OSHA's MDA rule, these methods will fall short of reliably quantifying dermal exposures. A newer technique, Video Imaging Technique for Assessing Exposures (VITAE) has been suggested as another alternative for dermal exposure assessment (Fenske, 1984). Given that current techniques are not developed enough to provide OSHA with a means of directly assessing and regulating exposures to MDA, a recognized dermal exposure problem, the introduction of another technique could be seen as a confounder in an already confused issue. Actually, this is an opportune time to explore options. Except for patch sampling to assess pesticide exposures, there are no institutionalized methods of assessing industrial, dermal exposures. Considering as many options as possible may prevent institutionalizing assessment techniques that do not optimally quantify exposures. Comparing the benefits and limitations of current methods and those of VITAE, VITAE (if validated) presents a tool to fill in many of the gaps left by current methods.

1.3 Overview of the VITAE Approach

VITAE is based on the principle that some chemicals fluoresce (emit visible light) when subjected to ultra-violet light. Some industrial chemicals, particularly some aromatic compounds fluoresce naturally (Fenske et al., 1986a). However, an industrial chemical that does not do so may be evaluated using VITAE by adding a tracer to the product at a known ratio to the chemical of interest. It is the presence of fluorescing compounds (tracer or target chemical) on skin surfaces that provide a marker that allows for the quantification of exposure.

After a worker has performed tasks in the workplace, body or clothing surfaces can be subjected to ultra-violet light and a video image taken. Fluorescent material on the surfaces will appear as bright areas on the image. By comparing the postexposure image with a preexposure image of the same surface, the amount of contaminant is determined by the change in total brightness of the image.

VITAE offers certain advantages over conventional approaches. First, VITAE is quick and noninvasive. It does not require application of solvents to the skin or hydration of the skin, as in handwash sampling, that might increase absorption (Bird, 1981). There is no need for laboratory analysis (after initial correlations are established) and VITAE does not require estimates of removal efficiencies. Additionally, the actual sample is retained as a permanent digital record and can be updated as analytical methods improve. Multiple samples taken throughout the day can be compared to baseline values to help pinpoint exposure episodes. Finally, the results provide a very graphic image that can be used as an invaluable

training tool for worker education.

For smaller operations, significant capital cost, not necessary for conventional methods, is required. However, in most processes that require ongoing monitoring of dermal exposures, the initial investment will probably be quickly offset by the costs laboratory analysis.

1.4 Statement of Problem

Although VITAE may be used without a tracer, if the contaminant of interest fluoresces strongly enough at the right wavelengths, in most cases a fluorescent tracer will be introduced to act as surrogate for the compound of interest. The amount of tracer necessary will vary, depending on the contaminant for which the tracer is a surrogate. For contaminants where it is necessary to measure very low surface densities (low target levels), relatively large amounts of tracer must be introduced. In some pesticide applications, a ratio of contaminant to tracer of as high as 5.7:1 has been used (Fenske, 1988). In other applications, where the presence of "inert" ingredients like the fluorescent tracers may dramatically impact the process or product, levels of tracer will need to be kept much lower. In fact, whether VITAE can be deemed a viable option will depend greatly on the amount of fluorescent tracer that can be introduced without degradation of the process and whether this amount of tracer is sufficient to be reliably measured by the imaging system. On the other hand, too much tracer can create problems for VITAE, regardless of whether or not it impacts the product or process. If the intensity of the light emitted by a high density of tracer on any portion of the skin exceeds the maximum level that the imaging system records, these high intensities will be recorded as intensity levels near the top of the dynamic range of the

imaging systems. Consequently, these greater than quantifiable intensities will be falsely recorded as lower levels and the method will underestimate the true exposure. Additionally, quenching may take place at higher surface densities of tracer that will result in unacceptable correlation between predicted and actual surface densities of tracer (Fenske et al., 1986b). Finally, use of the VITAE method by Black (1993) as part of an overall assessment of children's exposure to Chlorpyrifos from activity in treated lawns also revealed that variability of subjects' natural florescence (background brightness) can affect the intensity of florescence emitted by the tracer. Pervious work using VITAE sought to correct for background gray level by using a standard curve developed by exposing marked areas of skin with known amounts of tracer and developing a relationship between the background gray level and the irradiance of the tracer at different concentrations (Fenske et al., 1990; Black ,1993). However, the effectiveness of this method has not been determined, nor has it been determined whether using this approach of correcting for background gray level significantly impacts the outcome.

As these considerations indicate, an investigator wishing to use VITAE for dermal exposure assessment must introduce fluorescent tracer in amounts that fall within what can be termed an "operational window". The lower bound of this window is determined by the desired target surface density of the contaminant, the maximum amount of tracer that can be added without degrading the process, and the amount of fluorescent tracer that must be used to achieve surface densities of tracer that can be reliably quantified. The upper bound of the window is dictated by the upper quantifiable limit for the method. It becomes apparent that the investigator must know both the lower and the upper quantification limits for the VITAE

system before it can be determined if the introduction of an appropriate amount of tracer is feasible. These values will be impacted by several factors, including, masking of fluorescence from other materials present on the skin, interference from other fluorescing compounds, and variability in individual skin characteristics.

The preceding discussion presents several issues that must be addressed if VITAE is to be a valid tool for assessing occupational, dermal exposures. First, it must be demonstrated VITAE predictions correlate well to know tracer densities. Second, it must be shown that the impact of varying skin pigmentation (background grey level) can be controlled across the quantifiable range of the system. Third, a lower quantifiable limit must be established and exposures exceeding this limit reliably identified by the system. Finally, it must be show that an upper quantifiable limit can be determined, allowing investigators to calculate the maximum concentration of tracer that can be introduced and still produce acceptable correlations to contaminant concentration.

CHAPTER II

STUDY DESIGN

2.1 Introduction

A two phase approach was used to address the issues outlined in the preceding chapter. First, a summary standard curve was developed, using a wide range of skin pigmentations and doses so that the system is calibrated to handle a variety of conditions. From the images used for the summary curve, the parameters used to construct the curve and the assumptions made in using them were tested. How well the calibration data was described by the summary curve was evaluated by determining the tracer densities of the calibration images using the VITAE system and the known tracer surface densities. The second phase of the study was designed to test the method under simulated exposure conditions. Images were taken after applying two different exposure distributions to the hands of volunteers. The performance of the system when the distribution of surface contaminant was different from the distribution of the calibration exposures was evaluated using these images.

2.2 <u>Background of Summary Standard Curve</u>

As noted earlier, a summary standard curve that relates median background grey levels to tracer irradiance at different tracer densities has been used to adjust for the effects of background grey level. Variability of skin between individuals is, to say the least, complex. The amount of body hair, relative oil content of the skin, and skin imperfections like scars,

acne, or callouses are but a few of the myriad of factors that might impact the degree to which a tracer fluoresces. Attempting to identify, quantify, and correct for all variables would be too unwieldy. However, one gross measure of skin variability, background grey level (BGL), has been noted by previous investigators to noticeably impact the light emitting characteristics of fluorescent tracers (Fenske, et al., 1990; Black, 1993). The BGL of a skin surface may be determined by more than just the degree of skin pigmentation. Oil and water content, as well as body hair and other factors may influence the natural irradiance of the skin. Therefore, adjusting images based on a gross, descriptor like background grey level may actually incorporate the effects of other more specific factors.

The standard curve is, in effect, a model of relationships between natural background brightness, tracer surface density, and the change in brightness due to the addition of the tracer. None of the previous literature discusses the assumptions upon which the model is constructed or attempts to assess the performance of the method when the tracer surface distributions are different from those of the calibration method. To accomplish this, the modelling process must be detailed and the assumptions inherent in the model must be identified.

2.3 <u>Outline of Summary Curve Construction and Application</u>

The summary curve is constructed by using images of marked skin areas of human subjects. Images of the subjects are taken before and after varying, known amounts of tracer are applied in a consistent manner to the marked areas of skin (the process is detailed fully in the Methods chapter). Using VITAE software (Fenske, 1991) three histograms are produced; a preexposure, a postexposure, and a net histogram.

The preexposure and postexposure histograms provide the number of pixels at each of the 256 grey levels (0 - 255). From these histograms the median and average grey levels of the image and image brightness are determined. The brightness is simply the integrated area of the histogram.

A net histogram is generated by the VITAE software. It is produced by subtracting the number of pixels at each grey level in the preexposure histogram from those of the postexposure histogram and setting any negative values to zero. This means that the net histogram is not a true integrated difference between the postexposure and the preexposure histograms.

To develop the summary curve the median grey level of the pixels that represent exposed skin is needed. The postexposure histogram is a combination of two distributions; one of pixels that represents skin exposed to tracer and one of pixels that represents unexposed skin. Because of this, the median of the postexposure histogram is not an appropriate estimator of central tendency. Previous VITAE work used calibration images with high exposures and their net histograms to determine the number of pixels exposed to tracer in a calibration image (Fenske, 1990; Black 1993). Images of high tracer surface densities yield postexposure histograms where the distributions of the exposed and unexposed pixels are distinct. The net histogram, for these cases, were assumed to represent the pixels that correspond to exposed areas of skin. The median grey level of the net histogram for these images can be used for the summary curve because in this case the exposed pixel distribution can be separated from the unexposed pixels. All cases of bimodal postexposure histograms were identified and average number of pixels in the net histograms was

determined. Since the tracer was applied in a consistent manner to the same size area for all exposures, this average number of pixels was considered the number of exposed pixels and was used to determine the median grey level of exposed pixels for images where the two distributions of the post exposure histogram were not distinctly bimodal. This was done by using postexposure histogram. Starting at the highest grey level, the number of pixels for each grey level were incrementally added to a cumulative sum of pixels. The median grey level of the exposed pixels for distributions without distinct modes was assigned to the grey level whose contribution to the sum, made the cumulative sum equal to or greater than half the average number of pixels from the net histograms of the high surface density images. This grey level is referred to as the calibration grey level.

Images are then grouped according to their median BGL. Response curves are developed for each group of background grey levels (e.g. a single response curve might be developed for background grey levels 21 thru 23). The group response curves are linear regressions of the natural logarithm of calibration grey level of exposed pixels versus the logarithm of tracer skin loading (pg/pixel). The slope and intercepts of these BGL specific group response curves are then both regressed against the natural logarithm of the average of the median BGLs of each group. The resulting summary curve is actually two curves that describe a family of curves. The summary curve provides the slope and intercept of a linear curve that predicts how the BGL of a skin surface will impact the fluorescence of the tracer.

The VITAE software uses the summary curve information and the net histogram from image pairs of exposed subjects to predict the amount of tracer present on a skin surface. It does this grey level by grey level, applying the equation from the summary curve to each grey

level of the net histogram and then multiplying the result by the number of pixels in the grey level. This provides an estimation of the amount of tracer represented by each grey level. The sum of these provides an estimate of the total tracer present.

2.4 <u>Evaluation of the Model's Mathematical Relationships</u>

As discussed previously, the summary curve process is actually a method of modelling several imaging variables. If one considers a single pixel of an image from a tracer exposed skin surface, the brightness of the pixel will depend on the inherent brightness of the skin (background), the brightness due to the tracer loading, and a term that describes the interaction between the background and the tracer loading. This can be described conceptually in equation form as:

$$\ln Pixel\ Brightness = a \cdot \ln Background + b \cdot \ln Tracer + c \cdot f(\ln Background), \ln Tracer)$$
(Eq. 2.1)

Where

a, b, and c are constants.

The current summary curve method uses the median grey level of the preexposure histogram to describe the tracer surface density associated with each grey level in the net histogram. The description of the tracer density for a given net histogram grey level can be considered in equation form as given by:

$$\ln Mass / pixel = (C_1 \cdot \ln BGL + C_2) \ln Exposed + C_3 \ln BGL + C_4$$

(Eq. 2.2)

Where

 C_1 = Slope of Slope Summary Curve

 C_2 = Intercept of Slope Summary Curve

 C_3 = Slope of Intercept Summary Curve

 C_4 = Intercept of Intercept Summary Curve

Exposed = Individual Net Histogram Grey Level

BGL = Median Grey Level of the Preexpsoure Histogram

Equation 2.2 was derived from the empirical, graphic methods outlined by Black (1993). Although the VITAE calculating program uses several steps, it applies the same information in an equivalent manner to each grey level of the net histogram. This indicates that the surface density and the brightness are logarithmically related. Besides being supported by the empirical data, this is the recognized relationship between the optical density and brightness (Russ, 1992). However, the loading rate is not the dependent variable and the equation might be more appropriately written:

$$\ln Exposed = \frac{\ln Mass / pixel - C_3 \ln BGL - C_4}{C_1 \ln BGL + C_2}$$
(Eq. 2.3)

Equation 2.3 differs from Equation 2.1 and introduces several issues that may affect the performance of the system. First, as opposed to Equation 2.1, this equation is nonlinear (when using log transformed data). Also, there is no discrete term for the brightness that is due to the tracer density. Additionally, there is a peculiarity that exist due to the nature of the

equation. The equation breaks down when the average BGL is less than one. In fact, regardless of the surface density or the grey level of the histogram being evaluated, the calculated density approaches infinity as the average BGL approaches zero. This condition exists because the calibration method generates a negative number for C_3 . Since the grey level scale does not encompass absolute dark or bright, this may be a governing condition when the pigmentation of the imaged surface is very dark.

2.5 <u>Identification of System Parameters and Assumptions</u>

The VITAE uses the summary curve to describe how a specific element of skin, represented by a pixel, will fluoresce given the BGL of the skin element and the amount of tracer present on the surface. This interaction is modelled by exposing an area of skin several thousand times greater than the area represented by a pixel to a known density and measuring certain parameters that describe the distribution of pixels before and after exposure.

2.5.1 <u>Describing Distributions Using Measures of Central Tendency</u>

The summary curve method uses measures of central tendency to describe the distribution of pixels in the preexposure and net histograms. For the preexposure histogram, the method uses the median grey level. The calibration grey level described in Section 2.3 is used to describe the distribution of pixels that represent skin surface that is exposed to tracer. These exposed pixels are represented by the net histogram. The reason the median of the net histogram is not used is due to the way the method was developed. The calibration grey level predates the inclusion of the net histogram in the VITAE calculation program.

If these measures of central tendency describe the distribution well, they will be strongly correlated to the integrated brightness of their corresponding histogram. This was

tested by performing linear regressions of the central tendency parameters and their corresponding integrated brightness. In addition to a strong correlation, the intercept of the regression line should be zero. This hypothesis was also tested.

2.5.2 Symmetry of Distributions

The summary curve process treats calibration images as if the distribution of grey levels are uniform (all pixels have the same grey value). The preexposure images are assigned a single grey level (the median) and the exposed pixels of the postexposure image are treated as if the tracer density and resulting grey level are uniform when the response is related to the tracer density. Since the video ramp is linear, uniform distribution is not essential, but symmetry is. If the distribution is symmetrical, the pixels brighter than the central grey level will balance out the pixels darker than the grey level.

The output of the VITAE calculation program does not provide data that lends itself to any rigorous testing of symmetry for the histograms. However, if a linear regression of the median and the mean of the histograms yields a slope that is not significantly different than one, then the assumption of symmetry would not be unreasonable. In addition to this, the histograms were also visually inspected for obvious asymmetry.

2.5.3 <u>Independence of Samples</u>

Although it has not been addressed in any of the previously published literature that employs or evaluates the VITAE system, there are several assumptions about the independence of samples that are inherent in the summary construction. The construction of the summary curve requires that the images be grouped according to their BGL, but irrespective of either their anatomical location or the subject from which they were acquired.

This requires that the VITAE method be independent of these factors.

Fenske et al. (1990) noted that the response of the system did not appear to be subject or anatomical location dependent. However, these observations have not been supported with statistical testing. These assumptions of independence were tested by developing a response curve of the system using the same images that were used for the summary curve construction. If the response of the system is independent of both the subject and the anatomical location, so will the residuals of the response curve. These conditions were tested using the response curve described in the following subsection and analysis of covariance for the effects of anatomical location and subject.

2.6 System Response Curve

The predicted loading of the VITAE method can be considered an adjusted response of the system. It measures the response of the system and corrects for the effects of BGL. If the grey level range covered the spectrum of grey levels from absolute black to infinite brightness and if there was no electronic noise associated with the system, the response (i.e. the relationship between predicted and actual loadings) would be linear. However such is not the case.

The VITAE system is, in effect, a photon counter. The charge-coupled device (CCD) camera, imaging board, and filters employed in the system actually screen and count the photons that are associated with the irradiance of a surface of interest. The lower limit of the tracer density that can be reliably detected by this system is determined by several factors. First is the effects of electronic noise. In addition to the effects of temperature and power fluctuations on the performance of the more than 245,000 sensors that are associated with an

image, there is the simple fact that this many sensors cannot be manufactured to respond identically to the same input of photons. These factors create electronic noise in the system that, when photon flux is low, can be statistically important (Russ, 1992). The second factor that influences the density of tracer that can be detected by the system is the amount of tracer that is necessary to create enough change in brightness to cause a pixel in the image to raise at least one grey level. As noted earlier, this quantum requirement is dependent on the BGL. But there is a level at which all BGL will reveal a response. The third factor is experimental variability. Actual doses and areas are calculated from measured values and as such suffer from inherent variability. These factors are impossible to completely separate and determine the amount of tracer than can be detected above both the electronic noise, the quantum density requirements, and experimental variability. Together, they can be termed system noise and determine the extent of the low, flat portion of the theoretical response curve shown in Figure 2.1, as well as the transition to the linear portion of the curve. Surface densities were left in the log transformation since the system models the data to fit a linear curve in this form and evaluation of the models performance should be done in the same form.

The upper, flat portion of the curve, as well as the transition from the linear portion of the curve (Figure 2.1) is also influenced by several factors. The first, is suppression of tracer irradiance due to the effects quenching. The phenomena was noted by both Black (1993) and Fenske et al. (1990) and results from either tracer densities that are high enough to block UV radiation from reaching some of the tracer molecules and/or the presence of enough tracer that a significant portion of the tracer does not receive enough energy to fluoresce. The second factor is simple saturation of the detectors. The imaging board only

records grey levels to 255. Light intensities (photon flux) that would result in grey levels higher than 255, given the existing linear ramp, would be recorded as 255 (Rich et al., 1989).

As the system response gets higher, the contribution of system noise to the signal becomes less significant and the response curve should be related to irradiance of the imaged surface. This relationship should remain until quenching and detector saturation are significant contributions to the signal. If the images of the calibration doses are well described by the summary curve model, this portion of the curve, where neither electronic noise, quenching, nor detector saturation contribute significantly to the response, should lend itself to linear approximation. This model of system response, graphically portrayed in Figure 2.1, provides a way of testing how well the summary curve approach models the effects of loading and BGL, determining the operating range of the system, and testing the assumptions of sample independence outlined in the preceding subsection.

2.7 <u>Assumptions of Summary Standard Curve Application</u>

Another assumption of uniformity/symmetry is necessary for the VITAE program to apply the standard curve to net histograms. When the VITAE calculating program estimates the amount of tracer that corresponds to a given grey level in the net histogram, it treats the pixels in that grey level as if their distribution were uniform or symmetrical and as if their distribution were the same as the distribution of pixels in the preexposure histogram. This may not be true if features on the skin that tend to have abnormal grey levels (e.g. hair, scars, callouses, skin creases) also tend to either collect or avoid tracer differently than the

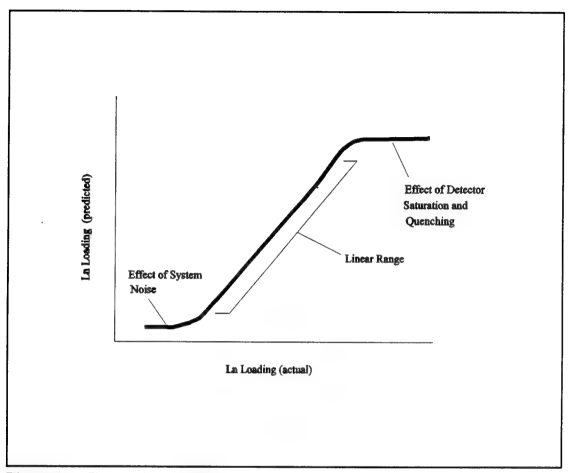


Figure 2.1. Theoretical Response Curve

remainder of the surface. Unfortunately, testing this assumption of symmetry is not feasible, given the imaging technology employed in the method.

The VITAE system uses the median BGL to construct the summary curve, but uses the mean BGL when it applies the summary curve data to each of the net histogram grey levels. This has the advantage of avoiding logarithms of zero for dark images with median BGL of zero. However, it demands that the two parameters do not differ significantly. If this is true, then the same test used to test the symmetry of the preexposure histogram will validate this assumption.

Probably the most critical assumptions that are associated with the summary curve application is the assumption of performance. If the summary curve is based on valid assumptions and relationships then it will correct properly for the BGL and the relative accuracy will not be dependent on the loading (density). This was tested using the regression of the In loading (predicted) vs the In Loading (actual). The residuals of the doses that fall along the linear portion of the curve were tested to establish that it was reasonable to assume their distribution was normal. These data were grouped according to dose (this will make the densities nearly equal). Analysis of variance was then performed to test the assumption that the model properly corrects for grey level. If the assumption is valid, then a significant portion of the variance of the residuals would not be described by the BGL. The performance of the system in predicting loading was tested in two parts. The first was to perform another analysis of variance, using the dose groups, to determine if a significant portion of the residual variance was described by the loading. The second part was to test the slope and intercept of the response curve for significant difference from an identity function. If the VITAE system predicts the loading perfectly, then the slope of the curve will be unity (one) and the intercept will be zero. These parameters were tested for the regression.

2.8 Operating Range of Summary Standard Curve

As with any analytical instrument, the operating range of the system must be determined to have a practical application. Black (1993) used changes in distribution to decide whether an image represented an exposed surface. This is a feature offered in the VITAE calculating program. The program detects changes on integrated brightness between the preexposure and the postexposure images and/or shifts in the maximum grey level. The

amount of change required to identify a surface as exposed is determined subjectively by the investigator. This approach has several drawbacks. First, the relative change in brightness is dependent on BGL. Therefore, a surface density identified by the criteria as exposed for one subject may be deemed unexposed for another subject. Secondly, the criteria have not been tested for any set of values and known exposures. A more straightforward approach was used for this study. The detection limit was calculated for the linear portion of the response curve. If the assumption of normality of residuals is correct and if the loading does not significantly contribute to residual variability, then the variance among dose groups should be relatively equal. Given this, the residuals can be pooled and the 95% confidence interval of the untransformed loading is given by:

95%
$$CI = e^{\ln Loading - 1.96 \cdot StdDev}$$
; $e^{\ln Loading \cdot 1.96 \cdot StdDev}$ (Eq. 2.4)

Where

StdDev = the standard deviation of the pooled response curve residuals

Equation 2.4 is not the 95% confidence interval of the line that describes the response curve. Rather, it is a confidence interval of the response of the system for any given loading (within the linear range of the model). Ideally, if the confidence interval can be expressed as a function of the true value, then setting the lower bound of the confidence interval equal to zero and solving for the corresponding loading would give the detection limit of the method. However, the model is based on equal variance across the linear range of the response curve and the response curve is a logarithm relationship. This implies that the variance of the

untransformed data approaches zero as the loading approaches zero. This may be more readily seen if the equation for the lower bound is rewritten as:

95% CI, Lower Bound =
$$\frac{e^{\ln Loading}}{e^{1.96 \cdot StdDev}}$$
(Eq. 2.5)

However, in this case the model also provides a practical solution. If the assumptions discussed earlier are met, the slope of both the model response curve and the curve of the untransformed data will be unity. Given this, the detection limit can be determined using the distance of the lower bound from the untransformed response curve. When this distance is equal to the loading, there will be greater than 95% confidence that these readings are associated with actual tracer response and not system noise. The detection limit then is given simply by:

Detection Limit =
$$e^{1.96 \cdot StdDev}$$
 (Eq. 2.6)

Equation 2.6 is the detection limit of the system and not the lower limit of quantification. The lower limit of quantification is also determined by the loading at which the response curve can be described as a linear relationship. This only becomes and issue if the detection limit falls below the lower end of the linear range.

As with the lower limit of quantification, the upper limit is driven by the linear portion of the response curve. The upper limit of detection will be determined using both the

assumption of unity for the response curve and the 95% confidence interval. If the addition of the samples from a dose cause a significant shift in the response curve from unity or if the average response of the dose falls outside the 95% confidence interval, the next lowest dose will be considered the upper limit of quantification for the method.

2.9 <u>Simulated Exposures</u>

Thus far attention has been focused on the summary curves, mainly determining if the assumptions inherent in their development and application are valid, how well the method models the calibration exposures, and what will be the range of the method. Even if the assumption are reasonable and the model describes the calibration data well, a critical question remains. Will the VITAE system respond comparably when the exposure conditions are not as constrained as the calibration exposures? In other words, it must be shown that the system response is not dependent on the distribution of the tracer on the skin. A practical way of testing this is to simulate exposures that might be expected in an occupational exposure, using known amounts of tracer.

Simulated exposures were done using the hands of volunteers. The hands were chosen for several reasons. They provide a convenient area to image, are easy to manipulate and do not require the removal of clothing. They also present rather complex surfaces, so they provide a good challenge for the system. Two exposure patterns were used, one involving the palm of the hand and the other using the dorsal portion of the hand. Palm doses were developed to simulate exposures that workers might receive when gripping contaminated surfaces. The exposures to the dorsal portion of the hand simulated deposition of droplets as might occur during spraying of pesticides or paints. Both exposures were

performed so that the amount of tracer deposited on the skin was known.

Images were analyzed using the VITAE system and the summary standard curve data. Response curves were constructed for both exposures by regressing the logarithm of the VITAE estimates of densities against the logarithm of the known tracer densities. The assumption that the system response is independent of distribution was tested by examining the parameters for the regressions of the two simulated exposures. If the assumption is valid, the regressed lines will not differ significantly from the response curve of the calibration images. That is, their slopes and intercepts will not differ significantly from the regression line associated with the calibration images.

2.10 Dose Ranges

Early work by Fenske (1986a) noted a detection limit of about 100 ng/cm², the beginnings of quenching effects at 700 ng/cm², and an upper quantifiable limit of about 2000 ng/cm². These limits were estimated from a gross examination of a standard curve produced using various tracer densities. The curve was developed without correcting for the effects of background grey level on tracer irradiance. Additionally, the study a used different camera, imaging board, and tracer. Black (1993), on the other hand, used the same equipment and tracer employed in this study. She noted a loss of linearity in the grey level response curves below a dosing concentration of 8 ppm and above 800 ppm. These limits translate to skin densities of approximately 12.5 ng/cm² and 1250 ng/cm². She provides no explanation on how it was determined linearity was lost or if an attempt was made to model the relationship between tracer density and irradiance outside these ranges.

Using the Turner 430 Spectrofluorometer, the lowest dose that can be reliably

administered to the palm of the hand is about 0.4 ng/cm². Dosing for the development of the standard curve and both simulated exposures thus began at this level encompassed a range that yielded surface densities of at least 1250 ng/cm².

CHAPTER III

METHODS

3.1 Equipment and Materials

3.1.1 Tracer

A fluorescent whitening agent (FWA) with the trade name Uvitex OB (2,2'-(2,5-Thiophene-diyl)-bis (5-tert-butylbenzoxazole) (Cas 7128-64-5)) was used as the tracer in this study. Uvitex OB was chosen as the tracer for the following reasons:

- 1. It does not exhibit significant toxicity (Review of toxicological data is provided in Appendix A).
- 2. The emitted fluorescence of Uvitex OB is fairly stable over time and under exposure to UV energy (Lee, 1990).
 - 3. Has adequate retention by the skin (solubility in water 0.01 g/100 mL).
- 4. Is readily soluble in acetone and toluene (0.5 and 5.3 g/100 mL respectively) to allow for dosing solution and proper elution from glass surfaces.
- 5. Uvitex OB has distinct and well separated extinction (375 nm) and fluorescence emission (435 nm) peaks.

3.1.2 Imaging Hardware

A. Camera:

All images were acquired using a Cohu 4810 monochrome Charge-coupled device (CCD). The camera collects 754 x 488 pixels.

B. Lens:

A Fujinon TV Zoom Lens - H6x12.5R was used for all image acquisition. The lens has eight principal f/stops (f/1.2 -f/16) and zooms from 12.5 to 75 mm.

C. Lens Filter:

A Kodak No. 2E Wratten Gelatin Filter was used to filter light entering the lens. The filter was secured to the lens end with a 52 mm technical filter holder. The filter blocks wavelengths below 410 nm, while transmitting light of longer wavelengths (~ 75%). This filter reduces interference from reflected UV and white light, but allows detection of fluorescence emitted by the Uvitex OB (435 nm).

D. Illumination:

Image surfaces were illuminated by two banks of four 4 foot F40 BLB (blacklight bulbs). The blacklights emit energy from 330 to 415 nm with the peak at 355 nm. The lights do not emit middle or short wave. Each bank of blacklights was filtered by selective glass filters developed by Fenske (1984). These filters further restrict the wavelengths illuminating the imaged surfaces, eliminating wavelengths above 400 nm.

E. Imaging Board:

Analog to digital and digital to analog conversions were accomplished with a Data Translation 2851 High Resolution Frame Grabber. The imaging board supported image capture (frame grabbing), image restoration, and computations. The imaging board provided a resolution of 512 x 480 pixels. This allows for the capture of less pixels (in both dimensions) than the camera senses. The excess pixels (upper rows and right columns) are

simply truncated with only the captured pixels appearing on the monitor screen. The imaging board assigns each pixel a numerical value to indicate relative brightness based on a 0 to 255 grey level scale. The scale, or video ramp is linear.

F. Computer:

Imaging software was run on an IBM-compatible computer (Dell 486P/33) with a 120 MB hard drive and 8 megabytes of RAM. Image manipulation (outlining and setting reference points) was performed using a Logitech three button mouse. Due to the memory intensive nature of image processing and limited hard drive space, images not being processed were stored on a 60 megabyte tape drive (Mountain Filesafe Tape Drive Series 7060).

G. Monitor:

Images were viewed for manipulation and examination using a Sony Trinitron Color Video Monitor (model PVM-1342Q).

3.1.3 Software

The software used to capture, display, and manipulate images, outline images, examine individual pixels, and provide pixel distribution data (histograms) was originally developed by Dr. William Gibb and was revised by Dr. Kyugon Cho. They are a collection of individual programs written in Microsoft C and utilizing software written specifically for the Data Translation video boards.

3.1.4 Analysis of Tracer Solutions

All tracer solutions were analyzed using a Turner 430 spectrofluorometer. Tracer was analyzed in solutions of toluene. The excitation wavelength of the spectrofluorometer was set at 355nm and the emissions were measured at 450nm.

3.2 Subject Selection

This study was unfunded. As such funds were limited and compensation of subjects to participate in the study was not available. Because of this, subjects were recruited from the students of the University of Washington's Environmental Health Program and their friends. Efforts were made to include as much ethnic diversity as possible, so that the full range of background grey levels and brightness that might be encountered in field evaluations would be included in this study. For the same reason, both males and females were included as test subjects. All subjects were briefed on the potential hazards of participation and the aims of the study. Consent forms were competed for each subject. A copy of the consent form is included as Appendix B.

3.3 <u>Imaging Technique</u>

The same imaging technique was used for both standard curve images and simulated exposure image sessions. There were differences in dose application, positioning of imaged surfaces, and image outlining in the two session types. Details for each are outlined in their corresponding sections later in this chapter. Following is a step by step overview of the imaging technique:

A. Setup:

The equipment was setup in a light tight room. The walls around the equipment were covered with darkroom curtains. The equipment was not moved during the study.

1. Camera: The camera was set level on a tripod with the lens center 100 cm above the floor. The f-stop was set on at 1.2 for all images. The focal length was set at 20 mm and the camera focused on the subject frame. Both the zoom and the focal rings were taped to

prevent movement when the lens was capped/uncapped or the f-stop was adjusted.

- 2. Subject Frame: The subject frame (constructed of 1 in by 2 in pine and painted flat black) was situated on a metal stand 125 cm from the detector of the camera. Its center was in line with the camera line of aim and 100 cm above the floor.
- 3. Lights: Each light bank was set on a stand with the center of the tubes oriented floor to ceiling and with their centers 100 cm above the floor. Both light banks were pointed toward the subject frame with the filter surfaces parallel to the subject frame. The distance between the light bank and subject frame planes was 90 cm. The centers of the banks were separated by 100 cm, making each center 50 cm for the camera line of aim.

The stands for the subject frame and light banks were connected at their bases to prevent inadvertent movement. Figure 3.1 provides a schematic view of the equipment layout.

B. Warm-up:

Prior to any imaging, lights and camera were allowed to warm up for at least one hour to allow the light emissions and camera system noise to stabilize.

C. Black Image:

Prior to all imaging sessions an image was taken with the f- stop closed and the lens cap in place. This image would be used later by the VITAE program to correct for system noise.

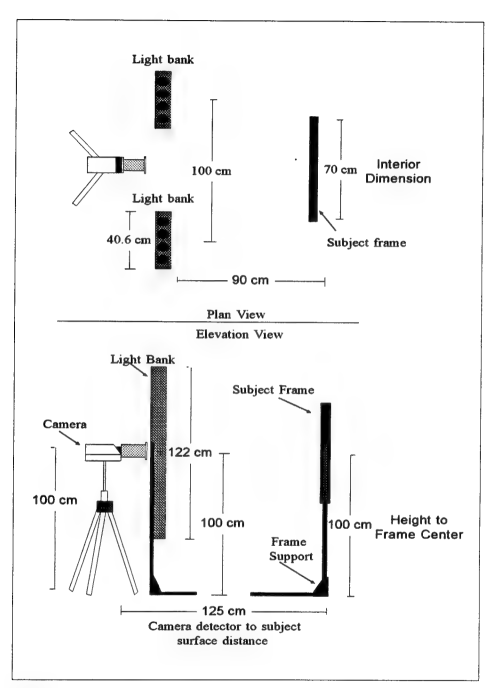


Figure 3.1. Equipment Layout

D. Standard Target - Initial:

An images of a standard target was used throughout the study to correct for fluctuations in the light energy produced by the black lights. An image of the standard target was acquired prior to the all subject imaging. Regardless of how many subjects were imaged in a day, each subject had separate standard target images. A piece of 75% cotton fiber white paper was used as the standard target. The same piece of paper was used throughout the study and care was taken to prevent soiling the target. The VITAE program prompts for initial standard target image as part of its image acquisition routine.

E. Preexposure Image:

Preexposure images were taken using the Image acquisition routine in the VITAE program. The standard body location identifiers in the program were not used, instead the "other" option was used for all image application. Awkward positions were avoided during imaging to make preexposure and postexposure images as comparable as possible.

F. Dose Application:

· Doses were applied as described in the pertinent sections of the following text.

G. Postexposure Image:

Postexposure images were taken less than three minutes after preexposure images. This was done to limit the possibility of crosscontamination or loss of tracer through unintentional contact of exposed skin surfaces. Since each subject had several doses applied and were required to reposition for imaging after dose application, subjects were carefully monitored and controlled to prevent inadvertent contact of exposed areas.

H. Standard Target - Session End:

Directly after the last postexposure image of each subject a session termination standard target image was taken. The same target used for the initial standard target image was used for the termination standard target.

3.4 <u>Summary Standard Curve Sample Collection</u>

3.4.1 Preparation of Doses

Ten exposure doses were made by dissolving Uvitex OB in acetone. The concentration of the doses were chosen so that lowest doses would yield surface densities below the linear range or the detection limit and the highest dose was above the concentration where quenching begins as discussed in Section 2.10. The interval between concentrations was chosen so that the natural logarithms of the doses were evenly spaced. This was done because the summary curve is based on the logarithms of the surface densities. Additionally, this spaces the concentrations closer together at the low end of the dose range, allowing for a more accurate determination of the detection limit. The doses developed were 0.28, 0.68, 1.44, 3.72, 9.52, 20.56, 61.12, 157.6, 425.8, and 1084 mg/L. As figure 3.2 shows, these doses do follow a log-linear increase from the lowest to highest dose. The same standard doses were used throughout experiment and were stored in capped 10 mL volumetric flasks, in 4°C freezer when not in use.

3.4.2 Special Setup

The equipment was set up in the same manner as described in section 3.3, except for the addition of a special target board. This board was constructed by painting a 72 cm x 72 cm x 1/4 in. polyfoam board flat black and cutting a 5 1/2 cm square hole in the center. The

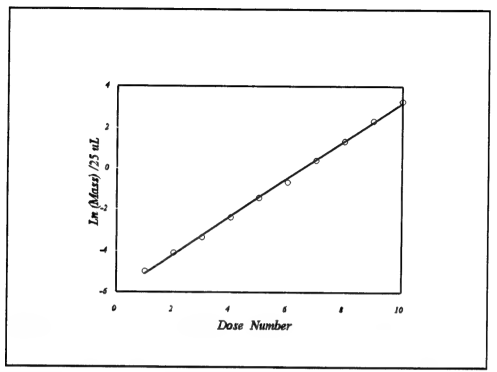


Figure 3.2. Distribution of Doses - Summary Standard Curve

board was attached to the back to the subject frame by four wing nuts, one in each corner.

3.4.3 Standard Target

The standard target image was taken by placing the target directly behind and flush to the polyfoam board.

3.4.4 <u>Target Selection and Preparation</u>

Targets (5 cm x 5 cm areas of subjects' skin) were marked using a 6 cm x 6 cm square of black construction paper with a concentric 5 cm x 5 cm hole cut in the center. Each interior corner was marked with a very small dab of fluorescent yellow paint. This was done to simplify the outlining process and to reduce error for any outlining where the boundary between the target marker and the subject's skin was difficult to visually distinguish. Target

markers were attached to areas of interest using a water soluble, nontoxic, pressure sensitive adhesive.

Eight areas were imaged for each subject. Areas were chosen from both the left and right palm, dorsal portion of the hand, inside of the forearm, and outside of the upper arm. Areas were chosen that presented a relatively planar surface. No surface preparation was performed on the imaged areas nor was effort taken to avoid or remove body hair.

3.4.5 <u>Preexposure Positioning and Imaging</u>

Preexposure images were not taken all at once but rather were taken as part of an image set. In other words, the first postexposure image was taken before the second preexposure image. Images were take by having the subject behind the polyfoam board with the marked area visible through the hole in the board. Since the hole in the target marker is slightly smaller than the hole in the polyfoam board there was some play in subject positioning. Prestudy trials revealed that making both marker and board holes the same size made it extremely difficult to ensure that the entire marked area was visible to the camera. A subject was considered properly positioned when all four fluorescent paint dots were visible on the monitor. Awkward positioning of the subject was avoided. This was generally done by having the subject find a comfortable position behind the board with the area of interest visible through the board hole and then placing the target marker in the same relative position.

3.4.6 <u>Dose Application</u>

Doses were applied in a controlled, random manner. Assignment of doses to each imaged area was performed prior to the imaging session. This was accomplished using a computerized random number generator, randomly assigning numbers from one to ten for

each of the eight areas. Doses for the palms and back of hands were controlled to ensure that at least two images for each dose were taken of both the back the hand and the palm. This was accomplished by rejecting an assigned dose if two images had already been acquired at the assigned dose for the controlled area (back of the hand or the palm). If a dose was rejected, another random dose was generated until it was not rejected. No distinction was made which side, right or left, had been imaged. This was done until two images at each of the ten doses were acquired for both the back and palm of the hand. Once these forty images were taken, any additional subjects imaged for the standard curve were assigned doses in an uncontrolled, random manner.

Doses were applied using a 25 uL positive displacement pipet (plunger - glass tube style). Using the pipet and 25 uL of the assigned concentration, the dose was applied by slowly depressing the plunger of the pipet while moving the tip across the marked area. Even distribution was obtained moving the pipet parallel to an edge, zigzagging back and forth down the length of the marked area and then repeating the process in a perpendicular direction. Care was taken to apply the dose about 1 cm from any marker edge and to apply the dose slowly enough that the solution would not wick outside the area of application.

For each summary curve image session four controls were taken for each dose used. Using the same pipet used for dose application, a 25 uL volume was placed in four separate sample jars. Toluene was added to each sample jar and the solution analyzed using the Turner 430 Spectrofluorometer. The mass applied for each dose was determined using these controls. This was done to ensure that no doses were contaminated during the study or that dose solutions were concentrating through evaporation from improperly sealed containers.

3.4.7 Postexposure Imaging

Postexposure images were taken as soon as the acetone was no longer visible on the surface. Since only 25 uL was used and it was spread of approximately 16 cm² this usually took less than a minute. The subject was positioned behind the board with the marker still in place. Again, it was ensured that all four florescent paint dabs were visible in the monitor before the image was acquired.

3.5 <u>Simulated Exposure Sample Collection</u>

3.5.1 Special Set-up

Images were acquired using the same image setup outlined in Section 3.3 except for the addition of two features to the image frame. First, a positioning line was added to the subject frame. The positioning line is made of black silk thread and was run horizontally across the image frame approximately 25 cm from the top. A small piece of marking tape (1 cm x 1 cm) was attached to the center of the positioning line to help subjects consistently position their hands. A shielding curtain was also added. The curtain, made of darkroom, cloth was hung from the top, back of the subject frame to cover the entire frame area.

3.5.2 Standard Images

For consistency, standard images for simulated exposure sessions were taken in the same manner as used for summary curve sessions. The black polyfoam board was in place, with the target paper directly behind.

3.5.3 <u>Preexposure Image Acquisition</u>

All images of simulated exposure were of both hands. Subjects were seated behind the picture frame with the middle finger of each hand touching the positioning line. This and the

marking tape, that was used for centering the hands in the subject frame, provided a way of consistently positioning the hands throughout the session. To ease the outlining process and to increase consistency, the hands were positioned with the fingers extended and pressed together. The subjects were asked to hold their hands so that the imaged surface was as planar as possible. The investigator ensured that the surface of interest (either the palms or the backs of the hands) were parallel to the subject frame. The shielding curtain was dropped between the hands and body of the subject so that the imaged surface was seen against a black background.

3.5.4 Dose Application

A. Palm Doses:

Doses were applied to the palm of the hands by having the subjects grip test tubes that were spiked with the tracer. Test tubes were spiked with solutions of acetone and Uvitex OB, using a 50 μ L positive placement micropipettor and the same method as described in Appendix C. Twenty spiked tubes were prepared for each subject (ten for either hand). The subject gripped each tube once, starting with the lowest dose and increasing in dose. Using the data develop in Appendix D, doses were applied to the tubes so that surface densities would begin below the anticipated detection limit (linear range) and exceed the density where quenching was anticipated. Exposure images were taken after the pair of tubes at each dose were gripped. For the first eight tube pairs, subjects were asked to grip the tube firmly with one hand lift it from the peg rack and return it to the peg rack. The contact time was approximately two seconds. The subjects were asked to contact the tube with the outside blade of the hand first to prevent deposition of tracer on the area between the index finger and

the thumb. Tracer deposited here would be missed when the palm was imaged. The process was repeated using the opposite hand and the other tube of the dose pair. The last two tubes for each hand were held lightly by the subject, while being slowly spun by the investigator. This was done in an effort to achieve detector saturation for some images.

B. Dorsal Doses:

Doses were applied to the dorsal portion of the hand using a 25 μ L positive displacement pipettor and solutions of acetone and Uvitex OB of varying concentrations. Each subject had six to eight doses to each hand, with the doses applied in increasing concentrations. Both the left and right hand were dosed and imaged together. Images were taken after each dose. The same dose was given to either hand at each dosing. The solution was applied by touching the pipettor to the skin surface, moving the pipettor form the surface, depressing the plunger so that a meniscus was developed extending from the end of the capillary tube, and then recontacting the surface. This was repeated until the tube was emptied and the plunger contacted the skin. The contacts were made in as random a manner as possible. To aid in this, doses were applied under white light where the tracer from previous contacts during the dosing and from previous dosing sessions would not be seen by the investigator.

3.5.5 Postexposure Image Acquisition

Postexposure images were taken as soon as the acetone on the skin had dried (about two minutes). Postexposure images were taken with the subject duplicating prexposure positioning as much as possible. Special attention was paid to ensure that the relative positions of the fingers and thumbs were constant.

3.6 <u>Image Pair Analysis</u>

3.6.1 Outlining

The VITAE system requires that the area of interest on the images be outlined. This is performed using a mouse. The program allows either straight line or freehand drawing, depending on mouse button manipulation. The postexposure image is outlined and then two reference points are chosen on the image. Next, the same two reference points were chosen on the prexposure image. From these, the program draws the outline around the prexposure image. If the fit is unsatisfactory, the reference points can be chosen again or the entire outlining process can be redone. The calibration images were outlined using the florescent paint dots. Straight lines were drawn using four points just inside the paint dots. The simulated exposures were outlined using a series of very short straight lines. This provided better control of the process. It was ensured that, whenever possible, no background (black area) was included in the outline.

3.6.2 Calculation of Tracer Mass

The VITAE system does not calculate the tracer density. It provides the mass of the tracer, which is a summation of the masses relating to each grey level. The program corrects for the effects of lens distortion, adjusting the histograms of the images before calculation of the mass associated with each grey level. On the other hand, corrections for variations in illumination are made when the total mass is calculated. The program uses ratio of the average standard target grey level to session end grey level to correct for difference in illumination between calibration and exposure measurement illuminations. The ratio of initial to session end grey levels are used to correct for variations in illumination within a session.

The VITAE program provides the mass in micrograms after the corrections have been made.

The density is calculated using this, the number of pixels in the outlined area, and the pixel dimensions determined in Appendix E.

CHAPTER IV

RESULTS AND DISCUSSION

4.1 Summary Curve Construction

4.1.1 Subjects

A. Results:

A total of 11 subjects were recruited for the summary curve portion of the study. The subjects yielded 88 image pairs, but only 82 pairs were used in summary curve development; one image pair was lost because the investigator failed to acquire a preexposure image, two images were discarded because postexposure images showed tracer solution had wicked beyond the marker boundary, and one image was lost in data transfer. Additionally, two images were not used because the preexposure brightness was greater than the postexposure brightness. The VITAE computer code generates a positive mass in these situations because after subtracting the preexposure histogram from the postexposure histogram, it sets all negative values to zero to produce the net histogram. Therefore, the magnitude, of what should be a negative response, can not be estimated. The demographics of the subjects are given in table 4.1.

B. Discussion:

The diversity of the subjects provided a wide range of median background grey levels.

The median background grey levels ranged from 0 to 51, which covers nearly one fifth of the dynamic range of the system as it was set up. This diversity in background grey levels allowed the method to be tested over a considerably greater range than was tested by

Table 4.1. Demographics of Summary Standard Curve Subjects

	Sex	
Race	Female	Male
African-American	0	1
Asian	1	1
Caucasian	2	5
Hispanic	ì	0

previous investigators. Using this calibration approach, Fenske et al. (1990) covered median background grey levels from 8 to 32, Black (1993) used levels from 3 to 23, and Archibald (1994) used grey levels from 4 to 18. Although these values are not directly comparable since the grey level measured for a surface is dependent on the lighting conditions, the equipment setup, and the equipment used, the range of grey levels as a fraction of the system dynamic range is comparable. This is due to the fact that all studies used the same imaging board, making the dynamic range equivalent, and the video response to light intensity is linear for the system. Given this relatively wide range of background grey levels, this study will provide the ability to test the system performance in correcting for the effects of background grey levels on tracer irradiance.

4.1.2 Grouping Images

As indicated by Fenske et al. (1990) the standard curve images were placed into groups based on the median background grey level of the image. The groups were chosen by trying to achieve groupings with similar sample size and without having images with the same background grey level in two different groups. Specific data for the images in each group are

supplied in Appendix F, while Table 4.2 provides a summary of grouping information.

Table 4.2 Summary Standard Curve Groups, by Median BGL

Group	N	Range of Median Grey Levels	Average Median Grey Level
1	12	0 - 9	3.17
2	12	10 - 14	12.67
3	12	15 - 17	15.92
4	11	18 - 21	18.27
5	12	22 - 29	25.87
6	12	30 - 38	33.25
7	11	40 - 51	43

4.1.3 Calibration Grey Levels

Of the 82 image pairs used, 19 had postexposure histograms that were clearly bimodal, with the modes well separated. This was determined through visual inspection of individual histograms. The average number of pixels in the net histogram for these images was 2426 (CV = 10%). Calibration grey levels were determined for each image pair using the 2426 pixel value, the postexposure histograms, and counting back method described in Section 2.3 of the preceding text. The value determined for each postexposure image is included in Appendix F.

4.1.4 Effects of Extreme Values on Linearity

The VITAE method requires that the logarithm calibration grey level be regressed (linear)

against the tracer loading (calculated in pg/pixel) for each background grey level grouping. As discussed earlier, Black (1993) noted a loss of linearity in this relationship, when the loading values were extreme. Because of the small sample size of each group and the variability present, rigorous methods of deciding when the impact of extreme loading rates affect the regression was difficult. Similarly, visual inspection of scatter plots had the same problems. The approach chosen was to perform several regressions on each group, each time deleting a group(s) of image pairs associated with doses at either end of the dose range. This approach is similar to that used by Fenske et al (1990). The effect of these deletions on the regressions is provided in Appendix G. Deletion of the lowest dose and then the two lowest doses affected the slopes and intercepts of the individual grey level group curves and the summary curves as well as resulting in a consistent increase in R2 for both the group and summary curves. Deletion of the highest dose, after the two lowest does have been deleted, does not make as dramatic an impact on the summary curve parameters and the effect on the correlation coefficients is inconsistent. For these reasons the summary curves were developed using the highest eight doses administered, but deleting the lowest two doses. The BGLspecific group response curves are presented as Figure 4.1 and the resulting summary curves are presented in Figure 4.2.

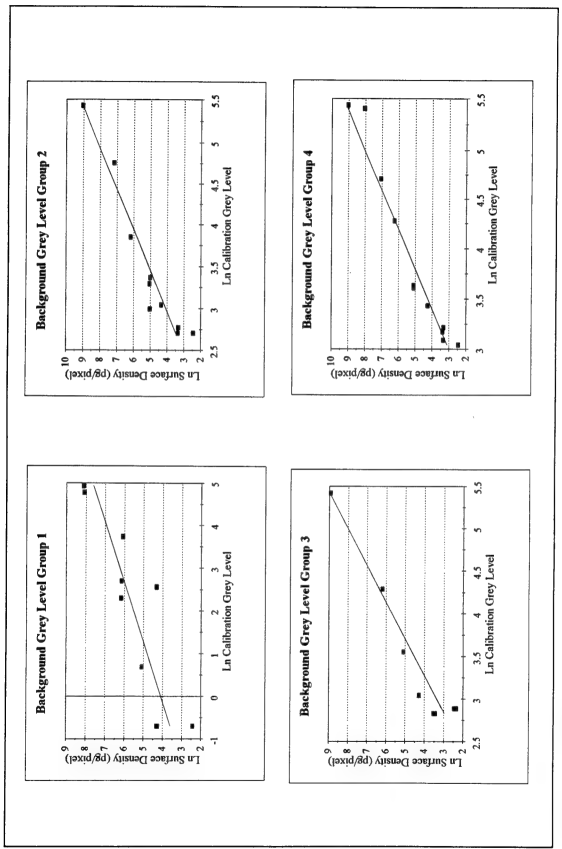


Figure 4.1 Background Grey Level Group Response Curves

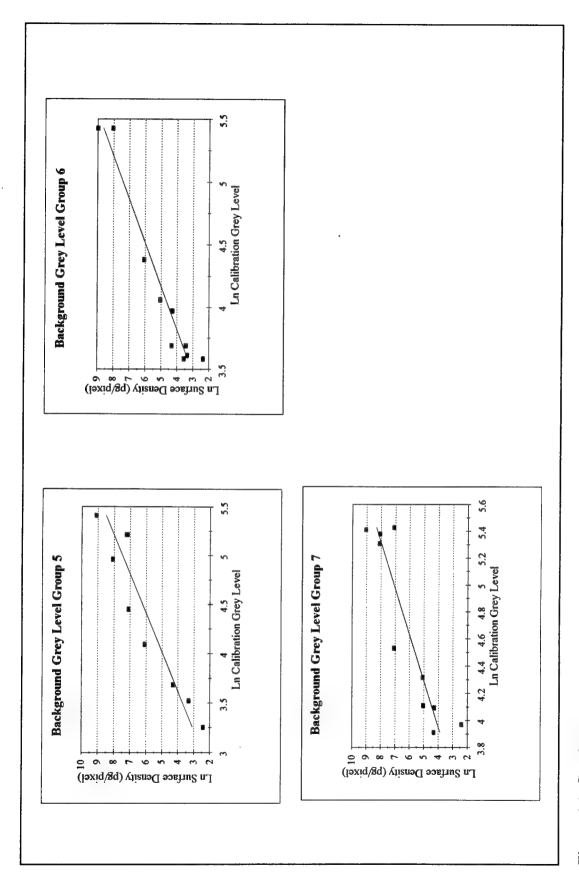


Figure 4.1 Continued

Figure 4.1 cont.

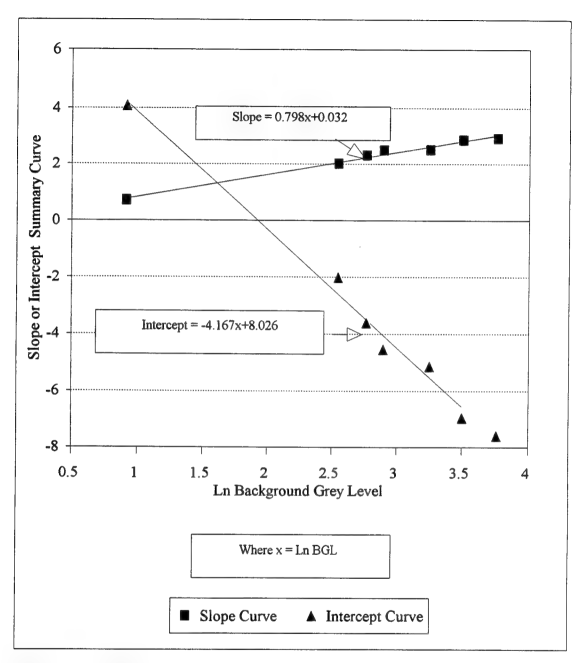


Figure 4.2. Summary Standard Curves

4.2 <u>Testing of Summary Standard Curve Development Assumptions</u>

4.2.1 <u>Describing Distribution Using Measures of Central Tendency</u>

As Figure 4.3 shows, the median of the preexposure histogram proved to be strongly correlated to the integrated brightness. A linear regression produced an R^2 of 0.978 and the constant (-2172) that was not significantly different than zero (p value = 0.15). The median grey level appears to be a good descriptor of the preexposure histogram.

On the other hand, the relationship between the calibration grey level and the net integrated brightness was not as good. Although the R² remained high (0.973), Figure 4.4 reveals this was probably due to the cluster of data points at either end of the regressed line. Despite the fact that the data can be described by a linear function (p value of the regression < 0.001), the data appears to actually be better described a nonlinear function. Additionally, the regression had a significant intercept (p value < 0.001). The intercept may be a manifestation of the way the net histogram is developed. By setting the grey levels with negative values to zero, after the preexposure histogram is subtracted from the postexposure histogram, the resulting net histogram may have an artificially inflated brightness. This inflation of the net brightness would contribute to the intercept term of the summary standard curve. The negative intercept term would shift the curve to the right, underestimating the exposure. An alternative is to use the average grey level of the net histogram instead of the calibration grey level to develop the standard curve. This would require no additional effort since the latest version of the VITAE calculating program generates this information. Given the way it is calculated, the function relating average net grey level and net brightness should not have an intercept.

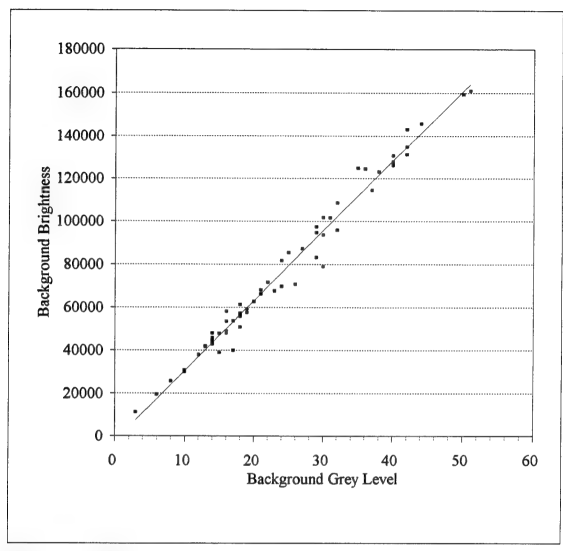


Figure 4.3 Background Grey Level vs. Background Brightness

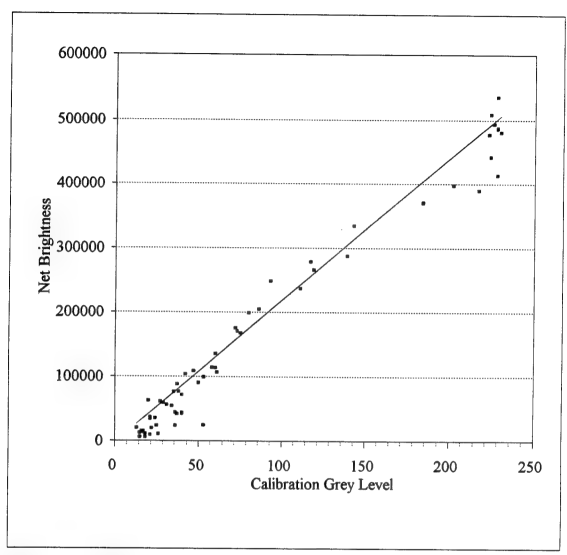


Figure 4.4. Calibration Grey Level vs Net Brightness

The use of the calibration grey level may have another impact on the system. Figure 4.4 reveals an interesting phenomena; there appears to be an upper bound to the relationship between the integrated net brightness and the calibration grey level. It appears that although the system measures a high response in terms of brightness, the calibration grey level does not capture this. Examination of high brightness histograms indicated that as brightness increases, the histogram seem to become left skewed. This shift in the shape of the distribution would cause a rise in the integrated brightness without a corresponding increase in a median-like parameter like the calibration grey level. What this means is that the quenching noted by Fenske et al. (1990) and Black (1993) might actually be partially due to distributional shifts and not the result of true, physical quenching of the tracer irradiance.

4.2.2 Symmetry of Preexposure Histogram

The linear regression of the average BGL and the median BGL (Figure 4.5) indicated both a very strong correlation (R^2 = 0.996) and a line that was not significantly different than unity. The slope of the regression line was 1.004, which, when tested for a slope equal to unity, could not be rejected (p value = 0.334). The same was true for a test of the intercept (0.295) not differing significantly from zero (p value = 0.283). These findings, coupled with visual inspection of the preexposure histogram, support the assumption that the histograms are reasonably symmetrical.

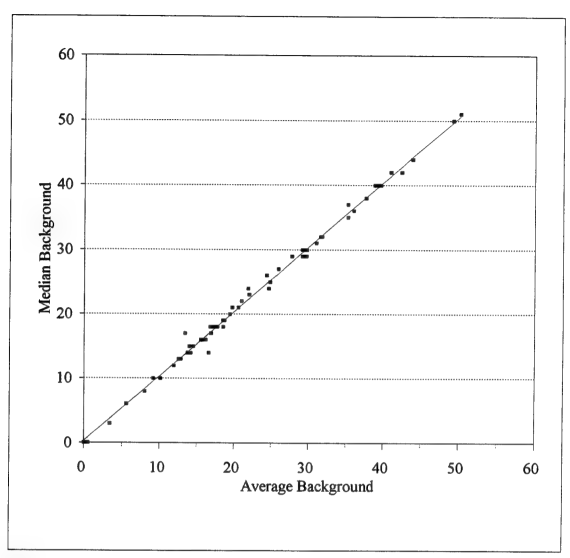


Figure 4.5. Average Versus Median BGL

4.3 System Response Curve

The system response curve is the linear regression of the logarithms of the actual loadings of the summary curve images versus the logarithms of the loadings predicted by the VITAE system. The regression line was generated using the doses identified in section 4.1.4 as being reasonably linear (doses 3 thru 10) and is provided as Figure 4.6. The two lowest doses that were not used to construct the summary curve are shown in Figure 4.6, but were not used in the regression of system response.

Initial examination of the response curve indicates that the summary curve approach models calibration exposures fairly well across the range of tracer densities used to develop the summary standard curves. The regression yielded a strong correlation ($R^2 = 0.945$). This is a rather impressive correlation given the fact that the loadings used in the regression spanned six orders of magnitude.

While the slope of the regression line (1.09) deviated less than 10% from unity, it was significantly different than one (p value = 0.01). Although the slope's departure from unity does not appear dramatic, the difference in the untransformed data will be much greater.

On the other hand, the intercept of the regression was rather large (-0.93) and significantly different from zero (p value < 0.001). This intercept is greater than 10% the linear range of the response curve, revealing a significant shift to the right. If the slope were one, this shift would translate to a systematic under estimation of tracer density of about 60%. However, the slope is not one. The identity function and system response curve converge. Table 4.3 illustrates the impact of the slope and intercept on the models prediction of tracer surface densities, given know tracer surface density.

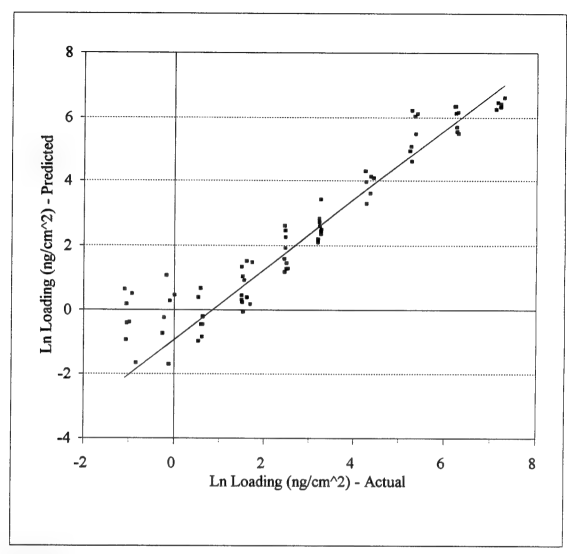


Figure 4.6. System Response Curve

Table 4.3. Examples of System Response for Untransformed Calibration Data

Surface Density (ng/cm²)	VITAE Predicted Density (ng/cm²)	Deviation from Actual Density (%)
2.8	1.21	-57.0
5	2.28	-54.4
10	4.85	-51.5
100	59.7	-40.3
1000	734	-26.5

Deletion of the highest dose did not create a significant change in the slope. However, this may be due more to the weight of the other data creating a stable line than the appropriateness of including this dose. All the data points associated with the highest dose fall below the regression line. It appears, looking at Figure 4.6, that quenching is becoming a factor at this loading. The fact that deleting this dose does not cause a significant shift in the slope may be an artifact of the method. The summary curve was constructed including this dose, so the model would reflect the influence of the images at the high dose.

Analyzing the calibration images also bore out the difficulties in evaluating images with average BGLs less than one. Six of the calibration images had average BGLs of less than one. The relative error of these images, for untransformed loadings, ranged from 1265 to 2.2×10^7 percent. The VITAE predicted loadings of these images were considered aberrant and were not included in any analysis involving predicted loadings.

4.4 Normality of Response Curve Residuals

Before the assumptions of independence for system parameters and system response can be evaluated, it must be shown that the assumption of normality for the residuals of the response is reasonable. Figure 4.7 provides a probability plot (probit scale). While not truly linear, it would appear reasonable to fit a line to the data. Using the residuals of the response curve and testing for normality with Lilliefors test, yield a Lilliefors' probability (2-tail) of 0.26. While the probability was not exceeding high, the assumption of normality was not rejected.

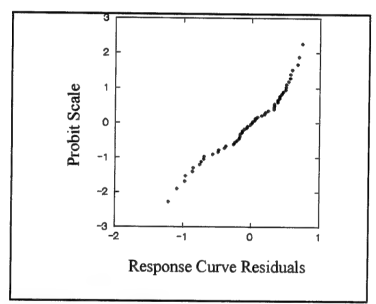


Figure 4.7. Probability Plot of Response Curve Residuals

4.5 <u>Independence of Samples</u>

Analysis of variance for the variables identified in subsection 2.5.3 indicate that the location of the sample does not contribute significantly to the variability of the system response. Evaluating the influence of anatomical location by using all eight locations as

different groups, yielded an R² of 0.127 and a p value of 0.432. If the dorsal portion of the hand, the forearm and the upper arm are consider similar surfaces, but dissimilar to the palm, grouping the areas in eight different categories might suppress a truly significant influence of anatomical location on system response. This was investigated by dividing the sample locations into just two group, palm and other surfaces. Analysis of variance on these categories reinforced the assumption that the system response is independent of anatomical location. This analysis yielded an R² of 0.019 with a p value of 0.304

On the other hand, subject differences described a significant portion of the variability, yielding an R² of 0.453 and a p value of less than 0.001. There is a question of valid analysis here. It would probably be more appropriate to perform the analysis with subject and BGL nested, since the variability might be more correctly attributed to skin pigmentation than some other factors of intersubject variability. Unfortunately, the statistical package employed could not perform the analysis on nested, dissimilar variables (one continuous and one categorical).

4.6 <u>Assumptions of Summary Curve Application</u>

As shown in subsection 4.2.2, applying the standard summary curve using the mean BGL, when the median BGL was used to develop the summary curve, should not create an invalid condition. The regression of median and mean BGLs yielded a line whose parameters (slope and intercept) did not differ significantly from an identity line (slope = 1, constant = 0).

The VITAE system also predicts the loading well across the linear range of the of the response curve. A linear regression of the loadings versus the residuals of the response curve validates that the performance of the system is not dependent on loading in this range. The regression produced a slope that was not significantly different from zero (p value = 0.96).

Analysis of variance for the effects of logarithm of the median BGL (this is the transform that was used to construct the model) on performance of the system revealed that the summary curve approach does not completely correct for the effects of background irradiance. The analysis yielded an R² of 0.80 (p value < 0.001). If the BGL was independent of any other variables, it would describe 80% of the variability of the response curve residuals. But two things should be considered when judging the affect of this seemingly dramatic correlation. First, the BGL is not independent of subject and a portion of the effect may be related to other factors of subject variability. Considering the fact that the correlation coefficient of the effect of BGL is considerably greater than that of effect of subject (0.45), it would appear that the reverse is probably true (i.e. variability described by subject is more correctly attributed to BGL). The second issue that should be considered is that this measure does not relate to relative performance. To answer this, the variability without the summary curve correction for BGL and/or the variability with some other approach must be known. This issue warrants investigation and should be a central issue in any further efforts to evaluate or enhance the accuracy of the system.

4.7 Operating Range of Summary Standard Curve

Before Equation 2.6 was applied to determine the detection limit an analysis of variance was performed to validate that the assumption that the response curve residuals were not dependent on the dose applied and could be pooled. This analysis produced a p value of 0.31, validating the assumption. The standard deviation of the pooled residuals was 0.531. Using this and Equation 2.6, the detection limit of the summary standard curve for the operating conditions of this study was determined to be 2.8 ng/cm². This corresponds to

loadings that would be produced by the application of a dose whose concentration lies between the third and the fourth dose. This falls within the linear range of the response curve, making it the lower limit of quantification for the system.

As noted earlier, deletion of the highest dose did not significantly impact the response curve. Additionally, the data points corresponding to the high dose did not fall outside the 95% confidence interval of the system response. The quantification limit of the system was not exceeded with the loadings used. But, as discussed earlier, this may be due to the method of developing the summary standard curve and not a lack of quenching.

4.8 <u>Simulated Exposures</u>

Eight subjects were exposed to tracer to evaluate of the system for performance under varying distribution conditions. All subjects were exposed to both simulated exposures in the same session. Unfortunately, data for three of the subjects were lost due to a tape drive malfunction. Again, image pairs with preexposure integrated brightness greater than postexposure brightness were deleted from the data set for the same reasons discussed earlier.

4.8.1 Evaluation of Dorsal Exposures

For each subject different concentrations of solutions were applied as described in the Methods chapter. Eight applications were made for each subject. The intent was to achieve loadings that started below the detection limit and exceeded the upper limit of the system. The regression for the response curve of the dorsal exposures was developed deleting images whose loadings were below the detection limit or had more than 5% of the pixels in the net histogram exhibiting saturation (having grey level of 255). However, this censored nearly half the data, leaving 48 of 80 image pairs. The remaining images only represented two of the

exposure concentrations used. To have enough data to allow for meaningful analysis, the lower doses were not deleted.

Figure 4.8 shows the response of the system as well as the regression of the logarithm of actual loading versus logarithm predicted loading, which is the response curve for the dorsal exposure. The regression line is described by a slope of 0.65 and an intercept of -0.045, both of which differ significantly (p values < 0.001) from the parameters of the calibration response curve. These regression parameters should not be considered reliable. This is due to the fact that the regression contains data below the detection limit of the system and the regression data is clustered in two distinct groups at either end of the data range. However, the data does suggest that under these conditions, the system responded differently than it did for the calibration exposures. This evident if only the exposures associated with the upper cluster are considered (they fall with the range densities for linear system response). In this cluster of surface densities, the predicted loadings for all five subjects and all exposures were below the actual loadings. The same is not true for predicted loadings of similar tracer densities for the system response curve. Additionally, the mean-square of the residuals increased more than three fold over the calibration response data, increasing from 0.318 to 1.396. This dramatic increase in variability is evident in Figure 4.8. The data points that fall well below the other data and the regression line, correspond to a subject whose BGL was considerably darker than the remaining subjects.

The distribution of the tracer was not as intended. The acetone solution wicked considerably when the skin was contacted with the pipettor. As a result, tracer was not distributed in a pattern of fine, round dots. Rather, it appeared, in the images, as small

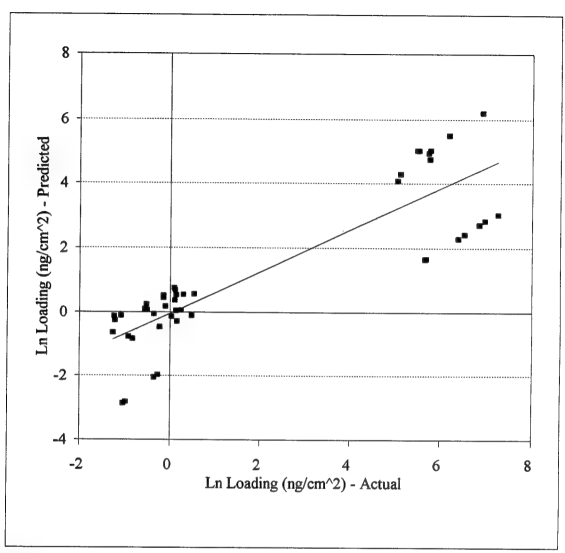


Figure 4.8. Response of Dorsal Exposures

asymmetrical patches. However, the distribution was still considerably different than the calibration exposures. Further, this may actually mimic true exposures more closely than the intended exposure pattern. Liquid aerosols deposited on the skin may wick similarly to the dorsal exposures of this study.

4.8.2 Evaluation of Palm Exposures

The simulated exposures to the palm of the hand were distributed across the range of loadings much more evenly than the dorsal exposure. This was due mainly to the method of application. The exposure pattern also appeared as intended, with relatively large areas of continuously exposed areas. The areas of exposure were the tip pads of the fingers and the portion of the palm just below the first finger joint. The brightness appeared nonuniform for exposed areas, with the intensity varying from low at the edges to high in the center of the exposed areas.

The distribution of doses allowed the response curve regression to be performed without using images with loadings below the detection limit or images with greater than 5% of the net histogram pixels exhibiting saturation. Figure 4.9 illustrates the resulting response curve. The lower doses (below a log transformed value of 1.0) are show in the figure, but not used in the regression. The slope of the palm exposure response curve was 0.718, which was significantly less than the slope of the calibration response curve (p value < 0.001). On the other hand, the intercept (-1.204) was not significantly different than the calibration response curve intercept (p value = 0.58). Although the response curve for the palm doses can be described with linear function (p value < 0.01), it appears in figure 4.9 that the data might be better described with a nonlinear function.

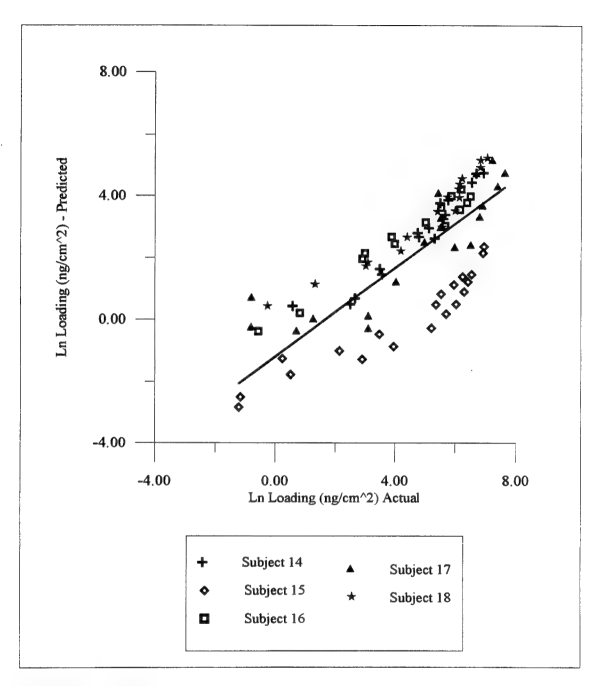


Figure 4.9. Response of Palm Exposures

Again, the logarithmic transformation of the data masks the true performance of the system. The theoretical (unity) response curve and the response curve of the palm exposures diverge. Using the regression line for the palm exposure response, the result of this divergence on the untransformed data is illustrated in Table 4.4.

Along with the large differences between actual and predicted tracer densities and increased variability, the simulated exposures of the palm also dramatically reveal the system's inability to completely correct for background irradiance. Figure 4.9 graphically illustrates this. Subject 15, who had skin complexion considerably darker than the other subjects, produced system response that was very different than the other subjects.

Table 4.4. Examples of System Response for Untransformed Palm Exposure Data

Surface Density (ng/cm²)	VITAE Predicted Density (ng/cm²)	Deviation from Actual Density (%)
2.8	0.63	-77.5
5	0.95	-80.9
10	1.57	-84.3
100	8.19	-91.8
1000	42.7	-95.7

4.9 <u>Potential Causes of Varying System Response</u>

The simulated exposures produced system responses that were considerably different than both the theoretical identity line and the response to the calibration exposures. The reason(s) for this varying response could not be identified with the approach used to evaluate the

system. But, the process of evaluation revealed several potential areas of concern that might contribute to the variations in response. These concerns might be due to either the construction and application of an inappropriate model or/and the effects of distributional differences.

The construction of the summary standard curves is equivalent to a rather complex mathematical model, providing the parameters of a linear curve for logarithmically transformed data. The parameters themselves are based on a functions that include multiple logarithmic transforms as well as untransformed values. The exponential of the equation (Equation 2.2) must be taken to obtain the estimated mass of the tracer. Given the relatively large values of the terms that determine the intercept and the fact that the intercept is negative for median BGL above two, there is the potential for the intercept to overwhelm any contribution from the slope term. Any problems associated with this might not be easily detected when the images evaluated by the system are similar (or the same) as the images used to construct the summary curve. However, when the distribution of tracer is different from the calibration images, the inappropriateness of the model may become apparent.

Another possible contributor to the variations in system response may be due not to the modelling approach, but to the differences in tracer fluorescence. Differences in fluorescence might arise from differences in tracer surface distribution. The calibration exposures were performed to achieve even, uninterrupted surface densities of tracer. As a result there was relatively little exposed/unexposed border length compared to the size of the exposed area. The same is not true for the simulated exposures. In these cases the exposed areas were not contiguous and ratio of border length to exposed area was greater than the calibration

exposures. Since the surface of the skin is not specular, a significant portion of the light emitted from a pixel sized area of skin and detected by the camera, might be light emitted from the area of interest and reflected off neighboring skin area to the camera. If neighboring skin areas are bright, as in skin exposed tracer, the neighboring surface would reflect more light than darker, unexposed skin. The result of such a phenomena would be reduced tracer irradiance for exposures with high exposed border length to exposed area ratios. Another difference between the distribution of tracer in the simulated and calibration exposure was the presence of an obvious density gradient. This may have resulted in a significant portion of pixels that contained insufficient amounts of tracer to be detected by the system.

CHAPTER V

CONCLUSIONS

The assumptions of central tendencies and symmetry inherent in construction of the summary standard curve were reasonable, with the exception of the calibration grey level. Despite the considerable amount of effort involved in determining the calibration grey level for each calibration image, the approach yielded a significant intercept when correlated to net histogram brightness. This, coupled with the fact that there appeared to be a upper limit to the brightness that could be represented by the calibration grey level, makes the parameter a less than optimal descriptor of irradiance of exposed pixels.

Optimizing the correlation coefficients of the BGL-specific response curves to determine the linear range the summary standard curve fails to reliably identify tracer densities that outside the linear range. The highest calibration exposure appeared to fall above the range of linear responses for the system, but optimizing the correlation coefficients of the individual BGL-specific response curves failed to identify this data set for censoring.

Although the system response curve had a high R² (.945), the response differed significantly from the theoretical identity curve. This may be due to the use of the calibration grey level and use of linear regression for possibly nonlinear data. The evaluation of the system response curve also revealed that the method is very sensitive to small changes in the parameters of the summary standard curves.

Evaluation of system response revealed that the response of the system is independent

of anatomical location of the imaged surface. In contrast, the summary curve approach does not completely correct for the effects of background brightness. Eighty percent of the variability of system response for calibration exposures could be described by the median BGL.

Evaluation of exposure to tracer with distributions that simulate some possible workplace exposures revealed that the VITAE method is not robust with respect to variations in tracer surface distributions. The method yielded estimates of tracer densities that were significantly different than the estimates obtained under calibration conditions. Additionally, there was a rise in the variability of the system response. The result was a dramatic underestimation of tracer densities, especially at the high end of the calibrated response range.

CHAPTER VI

RECOMMENDATIONS

The sole intent of the summary standard curve approach is to correct for the effects that background irradiance the skin on the irradiance of the fluorescent tracer. It is apparent of the method does not completely correct for this effect. However, it is still unknown as to what degree, if any, this effect is reduced. This should be determined before any further VITAE work using this approach is conducted. This could be accomplished by developing a calibration curve that does not correct for background irradiance (e.g. relating net histogram brightness to tracer mass) and comparing the results of such an approach to the results of the summary standard curve approach.

The effects of distribution on tracer irradiance should be explored and if the effect is determined to be significant, a means of adjusting for the effect must be sought. The effect might be explored by comparing the net brightness for several distinctly different tracer distributions with total tracer mass held constant. Artificial surfaces, where distinct boundaries can be developed and controlled, might also be used to explore the effects of boundary conditions.

An alternative for the calibration grey level should be sought to better describe exposed pixel distributions. One obvious candidate is the average net histogram grey level. Any alternative parameter can be tested comparing system response using the new candiate for describing exposed pixel distributions and response using the calibration grey level..

The distribution of pixels in the calibration image histograms (preexposure, postexposure,

and net) are all described by measures of central tendency. Use of a more complete description of the pixel distributions in the calibration approach may make the method more robust to changes in surface distributions of the tracer. Use of such parameters as integrated brightness, skewness, or peakness in different combinations should be explored.

GLOSSARY

<u>Brightness (integrated)</u> - The summation of all the grey levels in an image or portion of an image. From the histogram, this is calculated by:

Brightness -
$$\frac{255}{0}\sum$$
 grey level • # pixels at grey level

<u>Background</u> - Describes an image that represents a skin surface that has not been exposed to tracer. It is associated with some description of the image histogram (brightness or grey level). When associated with grey level, a term associated with the grey level's location in the histogram (median, mean, maximum, or minimum) precedes the use of the word background (e.g. median background grey level).

<u>Calibration Grey Level</u> - A median-like value that describes the distribution of the pixels that are considered to represent areas of skin that are exposed to tracer. It is calculated using calibration images where the distributions of pixels in the postexposure histogram are clearly bimodal, with one mode representing exposed pixels and the other unexposed pixels. The technique is more fully described in Section 2.3.

<u>Grey Level</u> - The digital value associated with a pixel that describes the intensity of light detected by the system. The hardware employed here gives a value from 0 to 255, where the increase in grey level corresponds to a linear increase in light intensity.

Histogram - calculated by the VITAE calculating program using the array of pixel location and grey level. It is the distribution (frequency) of grey level values in an image or portion of an image, represented in tabular form as the number of pixels for each possible grey level. (Rich et al., 1989)

<u>Image</u> - here is refers to the digital representation of the light intensities measured at the surface. The image is a 480 x 512 array. Each cell (pixel) in the array corresponds to a detector in the camera, which in turn corresponds to a finite area on the imaged surface. The analog equivalent is a picture element.

<u>Pixel</u> - is the cell in the image array. It has both two dimensional (height and width) and spectral (grey level) attributes. The number of pixels is set, determined by the hardware of the system. The spatial dimensions are a function of camera zoom and the distance from the camera to the surface imaged. Displayed in analog, it is the picture element of the image.

<u>Summary Standard Curve</u> - The VITAE calibration tool. It provides a linear relationship between the *In* of the postexposure grey level and the *In* of the tracer surface density, given the median background grey level. The summary standard curve is comprised of two lines, one describing the slope of the aforementioned relationship and the other describing the intercept. See section 2.3.

<u>Video Ramp</u> - Is the relationship between detected light intensity and assigned grey level. In

the case of this system, the video ramp is linear. If the grey level were plotted against light intensity, the resulting function would be linear.

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Appendix A

Review of Fluorescent Agent Toxicity - Uvitex OB

This review was taken and adapted from Fenske and Black (1990).

The fluorescent compound used as a tracer in this study belongs to a class of chemicals referred to as fluorescent whitening agents (FWAs). The FWAs (also called optical brighteners or fluorescent brightening agents) are widely used in textiles, detergents, papers, and plastics to achieve a bright white color. The FWAs absorb light in the invisible ultraviolet region and emit light in the visible (usually blue to blue-violet) region. The net effect is an addition of visible light to the substrate, making the substrate appear brighter (Kirk-Othmer, 1981). The FWAs generally have moderate to low acute toxicity. Many FWAs have been tested for chronic effects and do not appear to be mutagenic, carcinogenic, or teratogenic to experimental mammals. The compounds in general have not been shown to cause dermal irritation or sensitization in humans (Syracuse Research Corporation, 1979). of the estimated 45,000 metric tons of FWAs produced worldwide in 1975, 20,000 metric tons were used in detergents (Kirk-Othmer, 1981). FWAs are incorporated into detergents to adsorb onto clothing during laundering and maintain fabric whiteness.

Uvitex OB is not used as a detergent additive. Uvitex OB is the trade name for 2,2'-(2,5-Thiophenediyl)-bis(5-tert-butylbenzoxazole) (CAS 7128-64-5). This FWA is approved by the FDA as an optical brightener in food wrappers (21 CFR 178.3297(e)) and

is exempted from tolerances by the EPA, when applied to growing crops (40 CFR 180.1001(d)). It can be used as a fluorescent quality control agent for surfactants used in pesticide applications. Although these uses do not involve skin contact, the studies done to obtain federal approval for the uses provide a good toxicological database. The results of these studies were obtained under the Freedom of Information Act and reviewed.

The short-term studies revealed a rat oral LD₅₀ of greater than 10 g/kg (Ciba Limited, 1963; Thomann & Kruger, 1975). An accumulation study using rats reported no toxic symptoms, gross organ changes or changes in body weight gains when Uvitex OB was administered at doses of 0.025 and 0.25 mg/kg/day for 28 days. Tissues analyzed at the end of the study showed a negligible accumulation in the liver, brain, eyes, blood, fat of the testes, and fat storage tissue (Ciba-Geigy Limited, 1979). In one subacute oral study, rats were treated with concentrations of 50 ppm (2.5 mg/kg/day), 5000 ppm (250 mg/kg/day), and 50,000 (2,500 mg/kg/day) Uvitex OB in their diet for 14 weeks. After seven weeks of treatment, the high dose group showed no symptoms; the dose was increased to 100,000 ppm (5,000 mg/kg/day). No toxic symptoms or mortalities were reported in any group. No difference was found among the groups mean food consumption or body weight gains. Terminal urinalysis, hematology, gross and microscopic histopathology showed no treatment related differences. Differences in organ weights were observed among the treatment groups. The liver, kidney, adrenals, and ovaries appeared to be enlarged at various dose levels but, since no dose-response effect was observed, the differences were not considered to be related to the treatment (Ciba Limited, 1964a).

An oral study was also conducted with beagle dogs (Ciba Limited, 1964b). Dogs were fed concentrations of Uvitex OB in their diet equivalent to 2.5 mg/kg/day, 250 mg/kg/day, and 2,500 mg/kg/day for three months. No deaths were observed; no differences in weight gains or food consumption were noted. The only symptoms noted were loose stools in the intermediate and high dose groups. Terminal hematology showed no differences among the groups; terminal biochemistry and urinalysis showed scatter abnormalities in the dogs at various levels. No consistent abnormalities were observed in the microscopic histopathology. Organ weights varied considerably, but no difference could be attributed to treatment.

Effects of skin contact were tested both in rabbits and humans. In one study, 0.1 g of the material was placed in the conjunctival sac of the eye of six rabbits. The eyes of three rabbits were rinsed after 30 seconds. The eyes were examined at 1,2,3,4, and 7 days after exposure and irritation was noted (Ciba-Geigy Limited, 1975a). In another study, the sides and back of six rabbits were shaved and one side was scarified. Gauze patches soaked in Uvitex OB were applied to both sides and covered for 24 hours. No irritation was noted when the patches were removed or upon examination 48 hours later (Ciba-Geigy Limited, 1975b). Another study reports that application of technically pure Uvitex OB to rabbit skin resulted in minimal skin irritation (Thomann & Kruger, 1975).

In human irritation and sensitization studies, patches were soaked in 0.5% and 1.0% mixtures of Uvitex OB and soft white paraffin were applied to human subjects for 48 hours. No primary irritation was noted. A second application was made 2-3 weeks later for an additional 48 hours. Upon removal of the second patch, no sensitizing reactions

were noted. A total of 102 people were tested at both application strengths (Ciba Limited, 1964c). In another study, a 10% concentration of Uvitex OB in a detergent solution was tested in 64 human subjects. Nine applications were made over three weeks followed by a challenge application 2 weeks later. No subjects were sensitized (Griffith, 1973).

Two chronic studies were also conducted. In one study, 140 rats were fed Uvitex OB at a concentration of 1000 ppm in their diet for two years (Ciba Limited, 1968). No differences in growth performance, mortality rate, food consumption, terminal hematology, biochemistry, urinalysis, histology, and total and differential tumor incidence between the treated and control groups were recorded. Fluorescent material was noted in the body fat and eyes of the treated animals. Optic lens opacities in the treated animals were noted in excess incidence of that noted in the controls, although the numbers were small. Liver enlargement was noted in treated males although no evidence of altered histology, serum alkaline phosphatase activity, or serum glutamate pyruvate transaminase activity was observed.

A chronic study was conducted in which 104 mice were fed 1000 ppm Uvitex OB in their for one year (Ciba Limited, 1969). The mice were observed for an additional 26 weeks. Reproductive performance was observed after 36 weeks of treatment. Treatment continued through the F₁ generation. The progeny of the F₁ generation were reared to maturity under treatment, the sacrificed. No effect of treatment was observed in reproductive performance, litter parameters, or morphological abnormalities. Mortality rates, growth, food consumption, and tumor incidence were comparable between the

treatment and control groups. Fluorescent deposits were observed in the adipose tissue of the treated animals. A marginal increase in liver weight was observed in treated males.

Again no histological findings correlated with the increase weight. Four of the treated males had large liver tumors; if these were excluded from analysis, no significant difference in liver weights were found. The researchers concluded that there was no evidence that Uvitex OB may be carcinogenic.

Several estimates of skin contact to FWAs used a detergent additives have been made. One study attempted to measure the deposition of FWA on skin from using FWAcontaining detergents for dishwashing. Six subjects placed one hand in an FWA-detergent solution for 15 minutes, three times a day for six consecutive days. The average maximum deposition was 2 ug/cm² (Burg et al. 1977). Assuming a surface area of 500 cm² for both hands, this deposition results in a total deposition of 1 mg. Other estimates of adsorption of direct hand contact of detergent solutions are 0.1 mg on both hands (Buxtorf, 1975) and 0.07 to 0.17 mg (Gloxhuber & Bloching, 1979). A survey was done to determine the fluorescence on the hands of 104 housewives. Two housewives used FWA-containing detergents for dishwashing; they had an average of 0.03 ug/cm² (est. 0.015 mg total). The average fluorescence for those using FWA-containing detergents for laundry was equivalent to 0.106 ug/cm² (est. 0.053 mg total). Those not using FWA-containing detergents has a fluorescence equivalent of 0.086 ug/cm² (est. 0.043 mg total) (Burg et al., 1977). FWAs may also adsorb to the skin through contact with laundered clothing. One study found a transfer of 0.07 ug FWA/cm² from 48 hours of contact with a whitened fabric. Assuming a covered body surface of 1.5 m², a total of 1.05 mg would be deposited (Burg et al., 1977). Another estimate for transfer from whitened clothing is a range of 0.05 to 1.7 mg/day; although a range of 0.005 to 0.085 mg/day was thought to be more realistic (Buxtorf, 1975)> In contrast to these estimates, on survey of backs and feet found no fluorescence, indicating a lack of transfer from socks and shirts (Burg et al., 1977).

Appendix B Copy of Subject Consent Form

UNIVERSITY OF WASHINGTON

CONSENT FORM

INVESTIGATION OF SKIN DEPOSITION AND DETECTION PROPERTIES FOR FLUORESCENT WHITENING AGENTS

Investigator: Richard Fenske, PhD, MPH, Associate Professor Department of Environmental Health, 543-0916

Student Investigator: Keith M. Groth, Graduate Student Department of Environmental Health, 685-9299

24 HOUR EMERGENCY TELEPHONE NUMBER: 206-523-9799

PURPOSE

The purpose of this study is to evaluate an imaging technique developed to measure skin exposures to chemical substances which cannot be seen with natural light. This technique could be useful to workers who are exposed to chemicals and other substances in their work, e.g., agricultural workers, construction workers, laboratory workers, pharmacists. The technique uses a fluorescent compound known as Uvitex OB. This compound is widely used as a brightener in clothing, detergents and plastics (e.g., plastic food wrap). Although this compound cannot be seen under natural light, it can be seen under blacklight. Adding trace amounts of this fluorescent compound to chemicals that workers are exposed to, may provide a method to evaluate skin exposure to workplace chemicals.

There is no benefit of the subjects who participate in this research. The information gained may be of future benefit to society by providing a better method of measuring skin exposures to chemicals.

PROCEDURES

If you agree to participate, you will be asked to touch surfaces (glass tubes or plates) that have been treated with the fluorescent compound, Uvitex OB or a small amount of the compound dissolved in acetone. You will be asked to contact the surface briefly with your hands or several drops of the tracer/acetone mixture will be applied to your hands. Video images of the hands will be collected prior to and about 30 minutes after application of the tracer. The imaging procedure involves placing your hands under ultraviolet light, a blacklight, to illuminate the skin while taking photographs with a computer imaging system. These studies will take about 1 hour. You may participate in one to several sessions (up to about five). The number of sessions determined by the length of the study

and the mutual convenience of both you and the investigators. Volunteers will be asked to thoroughly wash their hands with soap and hot water directly after the session.

PHYSICAL RISK AND DISCOMFORT

The levels of acetone exposure in this study are very low and not expected to cause any harm. At much higher concentrations or prolonged exposures, side effects such as nose and throat irritations or skin irritations can occur. No risks are associated with exposure to the fluorescent tracer, since the compound is not considered toxic (harmful). The longwave ultraviolet light is not considered harmful, but in some cases may cause discomfort to the eyes. You will be provided with UV-A shielding glasses during image collection. Photographs made during the sessions will not identify any participant.

OTHER INFORMATION

Your identity will not be retained in the data for this study, and no records will be kept as to specific participants. The results of this study will be published in scientific literature, but only in summary form so that no individual data can be identified with any participant.

Participation is voluntary. You may choose not to participate and you may withdrawal from the study at any time without penalty or loss of benefits to which they are otherwise entitled.

Investigator's Signature ______ Date _____

SUBJECT'S STATEMENT

The study described above has been explained to me, and I voluntarily agree to participate in this activity. I have had an opportunity to ask questions and understand that further questions I may have about the research or about subject's rights will be answered by the investigator listed above.

Signature of the Subject		Date .	
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Copies to: Subject

Investigator's File

APPENDIX C

Glass Tube Elution Efficiency Study

Purpose

To determine the recovery efficiency of Uvitex OB from glass test tubes using toluene as the elution solvent.

Method

The following method of spiking and recovering tracer from glass test tubes was adapted from a method of recovering pesticides from test tubes developed by Fenske and Lu (1993).

Spike Technique

Test tubes (KIMAX, 127 mm x 16 mm i.d.) were spiked using a solution of Uvitex OB and acetone. All spikes were performed using 50 μ l of solution and a positive displacement micropipettor. The mass of Uvitex OB applied to the glass tube was varied by varying the concentration of the Uvitex OB/acetone solution. Uniform application was be accomplished by incrementally rotating the tube while releasing the pipet volume. The pipet was drawn back and forth in a zigzag pattern down the facing length of the tube from just under the label to about a centimeter short of the tip. Solution was released form the pipet at such a speed that the acetone evaporated without running. About one sixth of the pipet volume is released during application of the spike to this "face". The tube was rotated about one sixth of a turn and another sixth of the volume applied. This was repeated until the tube was evenly covered and the pipet volume was emptied. To achieve complete and consistent dispensing of the pipet volume, care was taken ensure that the plunger tip of the pipet was in contact with glass of the tube at the end of dispensing. Solution was applied below the label and avoiding the

tip. This resulted in a spike surface area of approximately 40 cm². All tubes were allowed to dry at least ten minutes before elution. Recovery efficiencies were determined for five concentrations that span the anticipated concentrations for doses delivered by contact with spiked test tubes.

Tube Elution

The test tubes were eluted by slowly pouring the toluene from a 30 ml tilting repeater dispenser over the spiked tube that was held about 30° from vertical. As the toluene was poured over the tube, the tube was slowly rotated counterclockwise one revolution and then rotated clockwise to the same start position. Tubes were eluted using either one volume of the dispenser (30 ml) or two volumes (60 ml). The eluate was collected in glass sample jars with foil lined lids. Samples were capped and shaken thoroughly. If samples were not analyzed immediately, they were stored in the sample jars in a freezer (normal temp.: -25 C). Control Samples

For each spiked concentration four control samples were made. Control samples were made by directly spiking the same volume and concentration that used for test tube application, directly into four separate samples jars. The same pipet used to spike the tubes was used for the control samples. Again, to achieve complete and consistent application, the plunger end of the micropipettor was touched to the inside glass of the sample jar at the end of the pipet volume discharge. Either 30 or 60 ml of toluene was added to the sample jar (depending on the eluant volume used for the samples) using the same tilting repeater dispenser that was used for the elutions. The controls were capped and shaken thoroughly. Controls were treated the same as their corresponding samples, stored or analyzed at the

same times.

Sample Analysis

Samples will be analyzed using a Turner Model 430 Spectrofluorometer. The spectrofluorometer was set with the excitation frequency at 355 nm and emission measured at 450 nm. The spectrofluorometer was calibrated immediately before sample analysis using standard solutions of Uvitex OB and toluene.

Results and Discussion

Table D.1 provides a summary of the elution efficiencies determined for various spike masses. Using two 30 mL elutions, as opposed to a single elution, did not significantly increase the elution efficiency for either the 28 mg/L, 50 uL spike or the 1196 mg/L, 50 uL spike (p-values > 0.2). But, two 30 mL elutions, instead of one, did significantly increase elution efficiency for the 197 mg/L, 50 uL spike (p-value < 0.05). Elution efficiency was not found to be a function of spike mass for either the single elution (p-value > 0.3) or the double elution. (p-value > 0.9).

Conclusions

Although using two elutions only increased efficiency for one spike mass, using two elutions consistently increased efficiency. For this reason as well as the fact that using two elutions should improve consistency since it will allow for recovery of any tracer that might have been missed due to a poorly performed first elution, two 30 mL elutions will be performed the for recovery of tracer form glass test tubes. The exception to this is when using two elutions creates the potential for the concentration of the elution to be less than the

Table D.1: Summary of Test Tube Elution Efficiency

Con. Spike	Sa	mple	S	Cor	itrols		Elution Volume	Recovery Efficiency	
(mg/L)	Avg. Mass (μg)	N	CV	Avg. Mass (μg)	N	CV	(mL)	(%)	
2.8	0.147	6	0.07	0.140	4	0.02	30	105.2	
28	1.44	6	0.02	1.41	3	0.02	60	102.5	
28	1.41	6	0.03	1.40	4	0.03	30	100.4	
197	10.17	6	0.02	9.87	4	0.01	60	103.1	
197	9.92	6	0.02	*			30	100.5	
1196	59.41	6	0.02	59.80	4	0.02	60	99.34	
1196	59.05	6	0.01	*			30	98.74	
1811	92.67	4	0.03	90.58	4	.01	60	104.61	

^{*} Used the values for the control with 60 ml eluant.

calibration range of the spectrofluorometer.

Expect for one spike mass, elution efficiencies were greater than 100%. This is possibly due to the loss of toluene to evaporation during the process and to the toluene remaining on the test tube due to surface tension. The average recovery efficiency for elutions using two 30 mL volumes was 102.4 %. Since this is near what appears to be typical the coefficient of variation for the different spike masses, no correction will be applied to the masses determined using this method.

Appendix D

Determination of Transfer Coefficient

Purpose

To determine the percentage of tracer transferred from a spiked glass tube to a subjects hand when the tube is gripped by the subject. The data was used to estimate doses delivered to the hands of subjects during simulated exposures.

Method

The same test tubes described in Appendix C were spiked in the same manner as described in the Appendix C. Subjects were asked to grip the test tubes, contacting the tube first with the outside blade of the hand. The spiked test tubes were resting upright on pegs and subjects were asked to establish a firm grip, lift the test tube from the peg, and return the tube to the peg. Total contact time was approximately two minutes. Six tubes were gripped consecutively with each hand. The tubes were then eluted as described in Appendix C and the elutant analyzed using the Turner 430 Spectrofluorometer.

Results and Conclusions

Table D.1 provides the results of the transfer coefficient study. No statistical analysis was performed on the data because only an estimate of the amount transferred was needed. It appears that the typical transfer coefficient is about 0.15. Transfer appears dependent on subject and not on spike mass or mass of tracer already transferred.

Table D.1. Results of Transfer Coefficient Study

Table D.I.	results of 1	ransier Coeificient	Study		
Subject	Hand	Spike Concentration (mg/L)	Spike Mass (ug)	Mass Remaining (ug)	Percent Transferred (%)
1	Left	4.12	0.215	0.149	30.48
1	Left	4.12	0.215	0.174	19.05
1	Left	4.12	0.215	0.164	23.81
1	Left	4.12	. 0.215	0.157	26.67
1	Left	4.12	0.215	0.178	17.14
1	Left	4.12	0.215	0.180	16.19
1	Right	4.12	0.215	0.168	21.90
1	Right	4.12	0.215	0.192	10.48
1	Right	4.12	0.215	0.205	4.67
1	Right	4.12	0.215	0.194	9.52
1	Right	4.12	0.215	0.180	16.19
1	Right	4.12	0.215	0.145	32.38
2	Left	1218	61.11	50.46	17.43
2	Left	1218	61.11	51.70	15.40
2	Left	1218	61.11	49.22	19.46
2	Left	1218	61.11	51.29	16.07
2	Left	1818	61.11	58.32	4.57
2	Left	1218	61.11	57.49	5.92

Table D.1. Continued

Subject	Hand	Spike Concentration (mg/L)	Spike Mass (ug)	Mass Remaining (ug)	Percent Transferred (%)
2	Right	1218	61.11	53.77	12.01
2	Right	1218	61.11	55.42	9.31
2	Right	1218	61.11	53.36	12.69
2	Right	1218	61.11	53.36	12.69
2 .	Right	1218	61.11	55.42	9.31
2	Right	1218	61.11	51.29	16.07
3	Left	1218	60.83	47.14	22.50
3	Left	1218	60.83	45.62	25.00
3	Left	1218	60.83	45.24	25.63
3	Left	1218	60.83	44.48	26.88
3	Left	1218	60.83	47.14	22.50
3	Left	1218	60.83	45.62	25.00
3	Right	1218	60.83	45.62	25.00
3	Right	1218	60.83	44.10	27.50
3	Right	1218	60.83	44.48	26.88
3	Right	1218	60.83	45.62	25.00
3	Right	1218	60.83	46.00	24.37
3	Right	1218	60.83	46.38	23.75

APPENDIX E

Determination of Pixel Dimensions at 20 mm Focal Length

Purpose

To determine both the vertical and horizontal dimensions that are represented as a single pixel on images taken by the VITAE system. This data will be used to approximate the size of areas of interest on digital images.

Method

The following method was adapted from a method used by Fenske (unpublished) The camera, lights, and subject frame were setup as outlined in Methods section, equipment layout subsection. An image was acquired of a 201 mm x 201mm white art board on a background of black darkroom curtain cloth. Using VTOOLS, pixels of the image were examined. The median grey level of the pixels of the white art board were estimated as were the median grey level of the black background area. The corners of the white art board were estimated by finding the pixel at each corner of the board that had a grey level closest to the average of the white and black area medians. The coordinates for each corner was recorded and the dimensions of each pixel found using the relative positions of each corner.

Results and Discussion

Determination of border corners

- Grey level of corners.
 - -- Median grey level of white art board: 9.
 - -- Median Grey level of black background: 184.

- -- Border corners selected for pixel nearest a grey level of 87.
- Coordinates of the corners of the white art board.
 - -- Top left: (59,154)
 - -- Top right: (60,384)
 - -- Lower left: (345,153)
 - -- Lower right: (347,384)

Determination of pixel area

- Calculation of pixel dimensions.
 - -- Horizontal target dimension (pixels):

$$L = ((384 - 154) + (383 - 153)/2 = 230 \text{ pixels}$$

Pixel Length = 201 mm/230 pixels = 0.87391 mm/pixel

-- Vertical target dimension (pixels)

$$H = (345 - 59) + (345 - 60)/2 = 285.5$$
 pixels

Pixel Height = 201 mm/286.5 pixels = 0.70403 mm/pixel

- Pixel area = $(0.87391)(0.70403) = 0.6153 \text{ mm}^2$
- H:L ratio = 0.6281/0.7791 = 0.8056

Conclusions

The manufacturer of the imaging board (Data Translation) list the pixel height to length ration as 5:6 or 0.8333. This is somewhat different that determined using the method outlined. However, Fenske (unpublished) determined a similar ratio (0.8062) using the same approach.

APPENDIX F Summary Standard Curve Data

	ber	els		15	o ₂	_	13	9	7	7	0	33	66	9	_
	Number	Pixels		1745	1320	461	1943	999	352	2407	2630	2603	2329	1400	211
	Net	Histogram	Brightness	29001	7210	2295	35195	1862	1235	266683	1749	104475	288387	20928	9865
	Median	Net	Hist.	51	5	4	81	7	3	120	9	41	139	51	12
	Mass/	pixel	(ng/pixel)	0.479	0.161	0.011	0.464	0.072	0.074	3.239	900.0	0.440	3.344	0.075	0.00
	Mapped	Area	(pixels)	3191	3187	3156	3293	3322	3199	3286	3029	3472	3183	3174	2872
	Calib.	Grey	Level	10	2	0	15	0	0	119	5	42	139	13	10
	Post	Exposure	Grey Level	4	0	0	9	0	0	92	4	36	92	11	10
- Group 1	Bckgrnd	Grey	Level	0	0	0	0	0	0	3	4	9	∞	∞	6
Table F.1. Summary Curve Data - G	Bckgrnd	Brightness		199	701	1154	1836	222	863	11344	11522	19635	25932	25679	22276
nary Cu	Tracer	Mass	(gn)	1.528	0.514	0.036	1.528	0.238	0.238	10.644	0.017	1.528	10.644	0.238	0 007
1. Sumi	Image			RUA	LHB	RHB	RFA	LUA	LFA	RHB	LHB	RHB	RHB	LHB	LHB
Table F.	Subject Image			11	11	11	11	11	11	10	10	-	4	4	_

	lber	Pixels		2250	405	2432	2003	903	864	1922	366	2747	494	1926	1450
	Number	Pix		22	4	24	20	6	8	19	3(27	46	19	14
	Net	Histogram	Brightness	61662	5903	278966	109354	12564	16113	63547	6137	486505	8955	59569	35104
		His	Bri	9	4,	27	10	1	1	9	9	48	8	5	3
	Median	Net	Hist.	28	15	117	54	<i>L</i> 1	18	33	11	228	18	32	24
	Mass/	pixel	(ng/pixel)	0.158	0.002	1.309	0.482	0.029	0.028	0.153	0.012	8.365	0.002	0.152	0.078
	N		(ng	0	0	1	0	0	0	0	0	8	0	0	0
	Mapped	Area	(pixels)	3247	3040	3010	3167	3248	3336	3365	3127	3240	3196	3382	3056
	Calib.	Grey	Level	27	12	117	47	15	16	20	15	228	16	29	21
	Post	Exposure	Grey Level	24	11	26	34	13	15	24	15	226	15	21	19
- Group 2	Bckgrnd	Grey	Level	10	10	10	12	13	13	14	14	14	14	14	14
- 1	Bckgrnd	Brightness		30196	30742	30763	37961	41742	41998	45881	435237	44679	43603	48098	42796
Table F.2. Summary Curve Data	Tracer	Mass	(gn)	0.514	0.007	3.940	1.528	0.093	0.093	0.514	0.036	27.102	0.007	0.514	0.238
2. Sum	Ітаде			LUA	RUA	LHB	LHB	LHB	RHB	LUA	LFA	LHB	RHB	RHB	LHB
Table F.	Subject Image			10	10	7	2	9	2	5	5	6	5	9	5

Table F.	3. Sum	mary Cu	Table F.3. Summary Curve Data - G	- Group 3							
Subject Image	Image	Tracer	Bckgrnd	Bckgrnd	Post	Calib.	Mapped	Mass/	Median	Net	Number
		Mass	Brightness	Grey	Exposure	Grey	Area	pixel	Net	Histogram	Pixels
		(gn)		Level	Grey Level	Level	(pixels)	(ng/pixel)	Hist.	Brightness	
2	LUA	0.017	49919	15	16	17	3447	0.005	61	12207	879
4	RFA	0.093	39117	15	17	17	2775	0.034	61	13815	706
4	LFA	0.007	36230	15	15	15	2613	0.003	15	2881	208
5	RUA	0.238	47778	15	19	21	3283	0.072	23	37573	1594
∞	LHB	0.036	48087	16	17	18	3077	0.012	20	11938	594
1	LUA	1.528	48998	16	62	73	3126	0.489	73	170691	2423
7	RUA	0.036	53618	16	17	18	3397	0.011	20	6252	320
œ	RHB	0.036	48423	16	17	18	3078	0.012	21	11737	557
9	LFA	27.102	58192	16	211	226	3585.5	7.559	226	493841	2784
10	RFA	0.093	39979	17	17	17	2975	0.031	20	15877	794
3	LHB	0.514	53778	17	30	35	3167	0.162	36	06191	2061
4	LUA	0.017	45881	17	17	18	3085	900.0	16	3714	226

Table F.	4. Sumi	mary Cu	Table F.4. Summary Curve Data - G	- Group 4							
Subject	Subject Image	Tracer	Bckgrnd	Bckgrnd	Post	Calib.	Mapped	Mass/	Median	Net	Number
		Mass	Brightness	Grey	Exposure	Grey	Area	pixel	Net	Histogram	Pixels
		(gn)		Level	Grey Level	Level	(pixels)	(ng/pixel)	Hist.	Brightness	
11	LHP	0.093	61473	18	21	22	3313	0.028	25	20326	807
1	LFA	1.528	50843	18	99	72	2942	0.519	70	175271	2537
1	RUA	0.514	55853	18	34	37	3138	0.164	37	88658	2326
2	LFA	3.940	57375	18	29	111	3408	1.156	111	237301	2165
80	LUA	0.514	59378	19	31	38	3162	0.163	42	77315	1774
8	RUA	0.093	59201	19	23	24	3162	0.029	26	35694	1343
9	RUA	0.036	57618	19	20	21	3089	0.012	23	0996	445
3	RHB	27.102	62816	20	222	230	3232	8.386	230	481197	2690
9	RFA	0.017	68864	20	20	21	3541	0.005	24	9167	405
2	RFA	0.093	66283	21	23	25	3352	0.028	28	24090	886
9	LUA	0.238	06089	21	28	31	3314	0.072	34	56978	1669
7	LUA	10.644	71669	22	218	223	3405	3.126	223	477397	2504

					T	П								
	Number	Pixels		380	2262	414	2212	2111	2207	1507	545	1552	1990	2497
	Net	Histogram	Brightness	9045	371273	11019	334871	135634	370479	54925	19732	71944	205241	508824
	Median	Net	Hist.	23	184	56	153	63	184	36	36	45	86	224
	Mass/	pixel	(ng/pixel)	0.007	1.284	0.011	3.218	0.445	1.355	0.028	0.007	0.073	1.200	9.046
	Mapped	Area	(pixels)	3267	3069	3188	3308	3436	2908	3340	3252	3252	3283	2996
	Calib.	Grey	Level	24	184	26	143	09	184	34	33	40	98	224
	Post	Exposure	Grey Level	23	137	25	105	48	148	32	31	35	90	224
- Group 5	Bckgrnd	Grey	Level	23	23	24	24	25	26	27	28	29	29	29
	Bckgrnd	Brightness		73254	67751	69902	81791	85548	70794	87373	100751	94941	97584	83317
mary Cu	Tracer	Mass	(gn)	0.007	3.940	0.036	10.644	1.528	3.940	0.093	0.007	0.238	3.940	27.102
5. Sumi	Image			RFA	RUA	LFA	RHP	LHP	RFA	LUA	LHP	RFA	RHP	LFA
Table F.5. Summary Curve Data	Subject Image			∞	7	∞	11	10	7	6	4	6	6	7

	Number	Pixels		2688	893	1059	1048	732	610	912	2087	2955	518	1672	1785
	Ź			<u> </u>											
	Net	Histogram	Brightness	488077	42783	44696	42463	30431	24115	44079	199022	536188	21543	99811	114966
	Median	Net	Hist.	228	48	41	40	41	39	48	68	228	41	59	63
·	Mass/	pixel	(ng/pixel)	3.075	0.076	0.035	0.028	0.002	0.011	0.031	0.432	7.842	900'0	0.073	0.157
	Mapped	Area	(pixels)	3462	3150	2649	3269	3366	3400	3002	3537	3456	2720	3249	3269
	Calib.	Grey	Level	228	40	36	37	38	36	40	08	228	38	53	28
	Post	Exposure	Grey Level	196	37	35	35	35	34	38	53	222	38	46	50
- Group 6	Bckgrnd	Grey	Level	30	30	30	31	32	32	32	35	36	36	37	38
Table F.6. Summary Curve Data - C	Bckgrnd	Brightness		102009	93829	79043	101804	110459	108598	95987	1240918	124624	97024	114691	123234
mary Cu	Tracer	Mass	(gn)	10.644	0.238	0.093	0.093	0.007	0.036	0.093	1.528	27.102	0.017	0.238	0.514
6. Sumi	Subject Image			RUA	RHP	LHP	LFA	RHP	LHP	RHP	RHP	LHP	RHP	LUA	LHP
Table F.	Subject			6	10	-	6	4	9	-	9	6	7	3	3

Number **Pixels** 2267 2169 1587 2193 2368 1788 1482 2321 915 2221 444 Histogram Brightness 114185 107252 413957 168140 398178 248763 91028 390215 442270 25293 44857 Mediar Hist. 228 100 217 224 202 55 76 49 57 64 (ng/pixel) 0.074 3.158 0.156 1.198 1.189 0.005 0.072 3.313 pixel 0.161 8.467 0.011 (pixels) Mapped 3315 3289 3217 Area 3184 3250 3288 3295 3230 3213 3201 3371 Level Calib. Grey 228 217 224 202 93 50 75 46 9 53 61 **Grey Level** Exposure 158 119 218 149 46 65 67 4 53 52 52 Bckgrnd Level Grey 40 40 40 40 42 42 42 42 44 51 Bckgrnd Brightness 126689 130809 134945 127940 126268 143140 134215 131303 159489 161083 145793 Tracer 10.644 3.940 3.940 0.238 0.514 27.103 0.514 0.036 10.644 Mass 0.017 0.238 (gn) Subject | Image | RUA RHP LHP RHP RHP LHP RFA LHP RHP LHP LFA 00 N 3 m ∞

Table F.7. Summary Curve Data - Group 7

APPENDIX G Effects of Extreme Values on Summary Curves

Table G.1 Effects of Extreme Doses on Linearity

Group or Summary Curve Parameter	N	Average Grey Level	Slope	Intercept	R ²
Using All Concentrations					
Group 1	12	3.17	0.704	3.421	0.374
Group 2	12	16.67	2.428	-3.75	0.714
Group 3	12	15.917	2.601	-4.885	0.87
Group 4	11	18.273	2.676	-5.273	0.933
Group 5	12	25.75	3.087	-7.989	0.862
Group 6	12	33.25	3.231	-8.778	0.838
Group 7	11	43	3.264	-9.272	0.821
Summary Intercept Curve			-5.061	9.154	0.991
Summary Slope Curve			1.019	-0.324	0.972
Deleting Lowest Dose					
Group 1	11	2.64	0.739	3.73	0.576
Group 2	10	12.8	2.034	-2.009	0.922
Group 3	11	16	2.5	-4.444	0.883
Group 4	10	18.273	2.511	-4.524	0.969
Group 5	10	25.8	2.519	-5.113	0.889
Group 6	11	33.364	3.023	-7.723	0.905
Group 7	11	43	3.264	-9.272	0.821
Summary Intercept Curve			-4.531	8.569	0.97
Summary Slope Curve	1		0.885	-0.124	0.975

Table G.1 Continued

Group or Summary Curve Parameter	N	Average Grey Level	Slope	Intercept	R ²
Deleting Two Lowest Doses					
Group 1	10	2.5	0.713	4.101	0.777
Group 2	10	12.8	2.034	-2.009	0.922
Group 3	9	15.9	2.321	-3.602	0.926
Group 4	10	18.1	2.511	-4.524	0.969
Group 5	10	25.8	2.519	-5.113	0.889
Group 6	10	33.1	2.869	-6.941	0.943
Group 7	10	43.1	2.933	-7.579	0.821
Summary Intercept Curve			-4.167	8.026	0.99
Summary Slope Curve			0.798	0.0332	0.981
Deleting Lowest Dose					
Group 1	12	3.17	0.704	3.421	0.374
Group 2	11	12.54	2.641	-4.387	0.718
Group 3	11	15.91	2.91	-5.806	0.722
Group 4	10	18.1	2.84	-5.825	0.883
Group 5	11	25.46	3.023	-7.754	0.838
Group 6	11	33	3.132	-8.408	0.746
Group 7	10	43.1	3.101	-8.609	0.784
Summary Intercept Curve t			-4.796	8.257	0.791
Summary Slope Curve			0.947	-0.062	0.882

Table G.1 Continued.

Group or Summary Curve Parameter	N	Average Grey Level	Slope	Intercept	R ²
Deleting Lowest and Highest Doses					
Group 1	11	2.64	0.739	3.73	0.576
Group 2	9	12.67 .	2.028	-2.02	0.846
Group 3	10	16	2.701	-5.05	0.723
Group 4	9	17.89	2.84	-5.82	0.883
Group 5	9	25.44	2.413	-4.7	0.875
Group 6	10	33.1	2.888	-7.21	0.845
Group 7	10	43.1	3.101	-8 .61	0.784
Summary Intercept Curve			-4.326	7.913	0.943
Summary Slope Curve			0.832	0.048	0.891
Deleting Two Lowest Doses and Highest Dose					
Group 1	10	2.5	0.7131	4.101	0.777
Group 2	9	12.67	2.028	-2.02	0.846
Group 3	8	16	2.382	-3.788	0.793
Group 4	9	19	2.583	-4.77	0.939
Group 5	9	25.44	2.413	-4.695	0.875
Group 6	9	32.78	2.705	-6.321	0.908
Group 7	9	43.22	2.732	-6.7634	0.967
Summary Intercept Curve			-3.912	0.753	0.982
Summary Slope Curve			0.7347	0.158	0.943

APPENDIX H Simulated Exposure Data

Table H.1. Simulated Exposure Data - Subject 14

			Actual		Pre	Pre	Pre	Post	Post	Post	Net	Ned	Net	Area	VITAE
Side	Hand		Mass (ug)	Pixels	Brightness	Median	Average	Brightness	Median	Average	Pixels	Brightness	Median	cm^2	Mass (ug)
Ъ	ч		0.080	17670	781871	47	44.259	17682	47	45.240	1522	74177	47	108.745	0.094
ď,	×	2	0.190	17200	751613	46	43.698	789168	47	45.561	2170	112887	55	106.525	0.164
а	~	3	1.532	17320	746346	46	43.092	804878	47	46.006	2508	134359	99	107.595	0.213
Ы	Я	4	3.851	17651	772430	46	43.761	861933	50	48.549	4286	251643	59	109.188	0.471
а	~	2	10.878	14789	670061	48	45.308	865925	58	58.366	6448	457600	99	91.242	1.328
Ь	Я	9	21.306	17271	771231	46	44.655	924369	50	53.559	3514	297470	76	106.144	1.469
Ы	~	7	27.847	15988	722004	47	45.159	1014331	58	63.171	7712	615187	69	98.751	2.925
D.	~	000	28.557	17449	772131	46	44.251	1028312	53	58.973	5121	453617	79	107.238	2.778
а	ĸ	6	70.572	16818	747871	46	44.468	1216935	56	72.389	7317	764382	83	103.389	8.634
ď	2	10	100.438	16311	717010	46	43.959	1265589	57	77.596	7135	828295	94	100.307	11.521
ሲ	u	-	0.080	17872	786183	45	43.990	768645	44	43.025	1522	57133	37	109.871	0.043
Д	L	2	0.190	17567	767016	45	43.662	759756	44	43.077	1803	67958	36	108.468	0.056
Ь	7	3	1.289	16939	745958	45	44.038	761868	45	44.564	2524	116298	37	105.141	0.171
Д	1	4	3.608	17948	780863	45	43.507	865990	47	48.220	2864	201859	70	110.449	0.566
a,	-1	2	10.635	16918	759219	46	44.876	905636	46	53.559	5001	391434	80	93.216	1.545
Д	ы	9	17.176	15160	682676	46	45.031	874519	53	57.697	4492	371488	79	103.991	2.013
P.	L	7	22.259	15224	808989	46	45.113	1011823	56	66.655	6104	580994	87	93.358	4.003
A.	L	∞	28.314	14701	669064	47	45.511	979118	53	66.725	4852	\$16341	101	90.246	4.257
Ы	٦	6	70.815	14563	692599	47	45.682	1162635	56	80.016	6071	753000	113	89.360	10.019
Д	L	10	129.838	16307	727448	46	44.610	1313263	54	80.533	5973	829068	126	100.289	13.894

Table H.1. Continued

					_					_					
VITAE	Mass (ug)	0.037	0.144	0.154	15.818	30.443	40.386	46.098	0.043	0.071	0.083	11.066	22.422	31.341	36.641
Area	cm^2	84.594	82.681	169.98	87.460	92.700	94.502	95.049	92.589	94.606	91.390	94.760	100.024	96.710	99.692
Net	Median	30	30	30	59	125	156	192	30	26	27	85	125	153	185
Net	Brightness	32879	122901	133183	890227	1409206	1748314	1909241	34942	62220	74071	677327	1127152	1487543	1606483
Net	Pixels	1092	3969	4333	8636	10096	11615	11543	1188	2284	2680	6237	8190	9776	9654
Post	Average	24.933	27.110	27.309	73.447	102.380	121.594	130.415	23.907	24.358	24.531	58.412	83.298	103.840	110.888
Post	Median	25	27	27	34	48	78	103	23	24	24	30	37	58	99
Post	Brightness	342959	364467	384947	1044491	1543170	1868428	2015568	359927	374694	364534	900015	1354762	1632878	1797487
Pre	Average	24.034	23.683	23.885	24.151	23.716	25.149	23.798	22.737	22.409	22.382	22.073	24.463	22.021	24.363
Pre	Median	24	24	24	24	24	24	24	22	22	22	22	22	22	22
Pre	Brightness	329654	318367	336975	343982	357569	383947	367917	342600	345417	331548	340494	398376	346780	395050
	Pixels	13716	13443	14108	14243	15077	15267	15460	15068	15414	14813	15426	16285	15748	16215
Actual	Mass (ug)	0.037	0.111	0.148	30.132	60.116	90.100	120.084	0.037	0.111	0.148	30.132	60.116	90.100	120.084
		-	3	4	S	9	7	∞	-	3	4	S	9	7	∞
	Hand	æ	æ	~	æ	æ	æ	ĸ	L	L	Ľ	L	L	ľ	L
	Side	Ω	Ω	Ω	Ω	D	Ω	Ω	D	Ω	Ω	D	Ω	Д	D
	_	_		_		_			_	_	_	_	_	_	

Table H.2. Simulated Exposure Data - Subject 15

Side Hund Actual Pre Pre Prof Prof <th< th=""><th></th><th></th><th>·</th><th></th><th>_</th><th></th><th>_</th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th></th<>			·		_		_															
Hand Actual Pre Pre Pres Prest Pres	VITAE	Mass	0.088	0.265	968:0	0.682	1.511	2.523	3.293	4.091	4.004	10.041	0.056	0.166	0.266	0.406	0.709	1.097	1.506	2.180	3.328	8.892
Hand Actual Pre Predian Average Brightness Median Average Brightness Median Average Brightness Pre Pr	Area	Sq cm	107.208	93.401	108.315	108.561	93.315	110.363	106.783	102.687	93.007	93.819	96.168	97.491	95.467	96.728	92.521	92.011	91.850	88.905	98.567	102.398
Hand Actual Pre Pre Pre Post Post Post Not Hand Actual Pixels Brightness Median Average 1536 45845 22 28.664 1538 <td< td=""><td>Net</td><td>Median</td><td>37</td><td>41</td><td>46</td><td>54</td><td>99</td><td>72</td><td>81</td><td>81</td><td>88</td><td>105</td><td>37</td><td>37</td><td>40</td><td>42</td><td>53</td><td>54</td><td>64</td><td>67</td><td>72</td><td>89</td></td<>	Net	Median	37	41	46	54	99	72	81	81	88	105	37	37	40	42	53	54	64	67	72	89
Hand Actual Pre Pre Pre Post Post Post Hand Mass(ug) Pixels Brightness Median Average Brightness Brightness	Net	Brightness	59537	151968	188693	214728	298589	420616	432226	506775	544064	766535	37586	103653	146319	166845	214895	266609	312029	377051	\$03509	728838
Hand Actual Pre Pre Pre Pre Pret P	Net	Pixels	1558	3584	4126	3754	4150	5347	4837	8699	5598	1099	1046	2754	3581	3788	3918	4500	4602	5163	6424	7000
Hand Actual Pre Pre Pre Pre Pre Prost Hand Mass (ug) Pixels Brightness Median Average Brightness R 1 0.034 17364 464852 27 26.391 499669 R 2 0.119 14853 398288 27 26.301 499669 R 3 0.927 17516 455439 27 26.301 484846 R 3 0.927 17586 462998 27 26.328 618430 R 4 3.489 17864 479966 27 26.328 618430 R 5 19.749 14920 419415 28 23.11 628829 R 6 27.748 17864 479966 27 26.386 784331 R 7 40.666 17349 461974 27 26.28 784351 R 8 52.066 15522	Post	Average	28.664	31.925	32.670	35.035	41.444	43.989	45.203	49.620	54.444	67.342	25.586	27.626	29.157	29.599	32.697	35.624	37.973	42.336	46.069	\$7.674
Hand Actual Fre Pre Pre Hand Mass (ug) Fixels Brightness Median Average R 1 0.034 17364 464852 27 26.590 R 2 0.119 14853 398258 27 26.5813 R 3 0.927 17516 455439 27 26.001 R 4 3.489 17586 462998 27 26.001 R 4 3.489 17586 462998 27 26.001 R 4 3.489 17586 462998 27 26.018 R 5 19.749 14920 27 26.328 R 6 27.748 17864 419946 27 26.628 R 7 40.606 17349 461974 27 26.628 R 9 63.407 15501 21533 27 26.038 L 1 <t< td=""><td>Post</td><td>Median</td><td>29</td><td>32</td><td>32</td><td>32</td><td>35</td><td>35</td><td>33</td><td>36</td><td>37</td><td>42</td><td>26</td><td>28</td><td>30</td><td>29</td><td>30</td><td>31</td><td>30</td><td>33</td><td>33</td><td>34</td></t<>	Post	Median	29	32	32	32	35	35	33	36	37	42	26	28	30	29	30	31	30	33	33	34
Hand Actual Pre Pre Hand Mass (ug) Pixels Brightness Median R 1 0.034 17364 464852 27 R 2 0.119 14853 398258 27 R 3 0.927 17516 455439 27 R 4 3.489 17586 462998 27 R 4 3.489 17586 462998 27 R 4 3.489 17586 462998 27 R 5 19.749 14920 419415 28 R 6 27.748 17864 479966 27 R 7 40.606 17349 461974 27 R 9 63.407 15071 415537 28 L 1 0.029 15522 415537 28 L 2 0.163 15534 376211 25 L <td>Post</td> <td>Brightness</td> <td>499669</td> <td>484846</td> <td>575380</td> <td>618430</td> <td>628829</td> <td>788383</td> <td>784851</td> <td>828505</td> <td>823359</td> <td>1027299</td> <td>40008</td> <td>437935</td> <td>452602</td> <td>465528</td> <td>491894</td> <td>532976</td> <td>567120</td> <td>612005</td> <td>738354</td> <td>960273</td>	Post	Brightness	499669	484846	575380	618430	628829	788383	784851	828505	823359	1027299	40008	437935	452602	465528	491894	532976	567120	612005	738354	960273
Hand Actual Pres Hand Mass (ug) Pixels Bnightness R 1 0.034 17364 464852 R 2 0.119 14853 398258 R 3 0.927 17516 455439 R 4 3.489 17586 462998 R 4 3.489 17386 462998 R 5 19.749 14920 419415 R 6 27.748 17349 461974 R 6 27.748 17349 461974 R 8 52.006 16706 435588 R 9 63.407 15071 411500 L 1 0.029 15322 415337 L 2 0.163 15324 3356114 L 3 1.748 15534 383630 L 4 5.039 15760 36414 L 5	Pre	Average	26.590	26.813	26.001	26.328	28.111	26.867	26.628	26.074	27.305	27.298	24.177	23.782	24.696	24.078	24.361	24.635	24.795	24.569	24.353	23.107
Hand Actual Pixels R 1 0.034 17364 R 2 0.119 14853 R 2 0.119 14853 R 3 0.927 17516 R 4 3.489 17586 R 4 3.489 17586 R 5 19.749 14920 R 5 19.749 17349 R 6 27.748 17349 R 7 40.606 17349 R 9 63.407 15071 R 9 63.407 15071 L 1 0.029 15384 L 3 1.748 15534 L 3 1.748 15534 L 4 5.039 15760 L 5 16.925 14906 L 6 27.354 14906 L 7 37.782 14861	Pre	Median	7.2	27	27	27	28	27	27	27	28	28	25	25	25	25	25	25	26	25	25	24
Hand Actual RR 1 0.034 RR 2 0.119 RR 3 0.927 RR 4 3.489 RR 5 19.749 RR 6 27.748 RR 10 96.189 LL 1 0.029 LL 2 0.163 LL 3 1.748 LL 5 16.925 LL 5 16.925 LL 6 27.354 LL 8 47.725 LL 8 47.725 LL 8 47.725 LL 9 60.097 LL 10 101.626	Pre	Brightness	464852	398258	455439	462998	419415	479966	461974	435588	411500	415537	376778	376211	383630	379472	366114	367217	368484	357133	387698	386068
Hand RR 1 RR 2 RR 3 RR 4 RR 7 RR 9 RR 9 RR 10 L 1 1 L 2 L 2 L 3 L 6 L 6 L 6 L 7 7 L 9 L 10 L 10		Pixels	17364	14853	17516	17586	14920	17864	17349	16706	15071	15222	15584	15839	15534	15760	15029	14906	14861	14536	15920	16708
Hand	Actual	Mass (ng)	0.034	0.119	0.927	3,489	19.749	27.748	40.606	52.006	63.407	96.189	0.029	0.163	1.748	5.039	16.925	27.354	37.782	47.725	60.097	101.626
			-	2	3	4	5	9	7	00	6	10	-	2	9	4	'n	9	7	00	6	10
Side		Hand	2	~	×	×	×	ĸ	Я	×	×	ĸ	L	Г	L	L	1	Г	L	L	L	L
		Side	а	Ы	ы	Д	а	Ь	d,	а	Q,	Д	Ь	Ь	М	a.	а	Ы	Ы	Д	Ы	ď

Table H.2. Continued

			Actual		Pre	Pre	Pre	Post	Post	Post	Net	Net	Net	Area	VITAE
Side	Hand		Mass (ug)	Pixels	Brightness	Median	Average	Brightness	Median	Average	Pixels	Brightness	Median	Sqcm	Mass
D	R	-	0.037	16873	128049	9	7.589	130608	9	7.746	1816	18119	•	103.702	0.059
Ω	М	2	0.074	17091	129725	9	7.590	137848	9	8.070	1911	23065	10	105.055	0.136
D	R	3	30.058	16978	128271	9	7.555	486833	6	28.666	5786	389711	46	104.446	5.460
D	R	4	60.042	16304	124184	9	7.617	810551	14	49.523	7417	723893	85	100.658	9:636
Ω	×	5	90.036	15495	105007	9	6.777	990082	20	63.913	8913	928711	6	95.270	14.527
Ω	×	9	120.01	17292	130279	9	7.534	1270146	28	73.453	9738	1189879	124	106.347	17.595
Ω	L	-	0.037	16052	107840	9	6.718	110190	9	6.868	665	8741	6	98.665	0.059
Ω	L	2	0.074	15649	104708	9	6.691	122047	9	7.762	1888	25152	10	769.96	0.136
Ω	L	3	30.058	16624	109657	9	6.596	480504	10	28.899	6654	400316	41	102.257	5.460
D	L	4	60.042	14318	81659	5	5.700	671808	12	46.917	7751	629532	71	88.063	9:636
Ω	L	S	90.026	13738	75826	5	5.519	908614	35	66.129	9222	877718	91	84.502	14.527
D	L	9	120.01	13709	75198	5	5.485	1053023	51	76.823	9833	1026125	104	84.299	17.595

Table H.3. Simulated Exposures - Subject 16

VITAE	. Mass (ug)	0.103	0.153	1.098	1.910	3.006	4.788	6.864	8.173	19.917	0.088	0.206	0.938	1.505	2.637	4.384	5.442	6.446	17.196
Area	(sq cm)	133.898	125.023	128.756	131.592	129.827	129.470	126.499	121.991	131.321	127.662	111.782	130.700	128.178	127.551	126.161	124.581	120.165	123.597
Net	Median	45	54	64	89	69	82	98	93	111	51	58	19	64	99	80	88	96	102
Net	Brightness	101184	109258	468369	627400	684072	680235	787340	809130	1130218	72421	126321	441568	569002	609121	620869	688602	694857	1048160
Nat	Pixels	2191	2054	7020	8480	8379	7245	7744	7430	8451	1527	2214	7011	8318	1760	7305	7017	6549	8315
Post	Average	45.024	47.240	54.809	59.219	62.177	64.182	68.623	71.577	82.134	43.631	46.552	51.301	55.137	58.291	61.856	63.814	65.908	79.050
Post	Median	46	49	55	58	58	55	57	57	58	45	48	52	53	53	52	52	52	54
Post	Brightness	982281	968179	1151102	1275811	1310570	1352113	1417278	1419587	1760635	905958	8511620	1088197	1153641	1208265	1269419	1293908	1286592	1585660
Pre	Average	45.208	45.984	44.958	45.173	45.361	45.293	45.318	45.601	45.145	43.036	44.288	42.663	42.641	43.158	42.755	43.006	42.525	42.559
Pre	Median	47	47	47	47	47	47	47	47	47	45	46	45	45	45	45	45	45	45
Pre	Brightness	984276	934818	941233	966562	9\$7576	953524	932155	904549	963991	893346	804981	906673	888717	895107	877077	871174	830894	855305
	Pixels	21772	20329	20936	21397	21110	21052	20569	19836	21353	20758	18176	21252	20842	20740	20514	20257	19539	20097
Actual	Mass (ug)	0.073	0.284	2.555	6.347	19.359	31.879	43.908	58.393	168.373	0.073	0.318	2.393	6.873	36.086	57.935	72.419	78.557	170.862
		-	2	3	4	S	9	7	00	91	-	7	3	4	S	9	7	00	10
	Hand	Я	ĸ	×	ĸ	Ж	R	ĸ	×	×	ľ	ľ	ľ	L	ľ	IJ	Г	Г	L
	Side	Ь	Ъ	а	Д	Ь	Ы	Ь	Ь	Ч	Ь	Д	Ы	Ы	Ъ	Ъ	Ы	ፈ	Ч

Table H.3. Continued

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VITAE	Mass (ug)	0.107	0.126	0.190	0.251	22.978	37.368	47.928	52.355	0.065	0.135	0.207	0.272	18.532	29.827	34.210	40.643
Area	(sq cm)	120.546	116.604	121.358	124.962	125.681	130.466	128.547	133.443	123.228	122.514	121.155	127.200	120.891	120.595	125.122	118.406
Net	Median	31	32	33	33	73	150	190	211	27	30	32	31	72	160	141	185
Net	Brightness	90593	116504	160975	210562	1174163	1638173	1984430	2131501	50196	101097	148469	192860	1091541	1456276	1657115	1829549
Net	Pixels	2838	3608	4772	6242	10033	11084	12203	12344	6691	3081	4372	5773	2686	9704	11584	11384
Post	Average	27.426	28.264	28.973	29.560	72.210	91.298	107.069	110.833	24.465	26.184	27.475	28.131	68.868	87.746	93.523	106.068
Post	Median	27	28	28	29	35	37	43	46	24	26	27	27	35	37	42	47
Post	Brightness	537252	535950	571811	600677	1474891	1932773	2244922	2405408	490113	521555	540936	582416	1353320	1720532	1902162	2042240
Pre	Average	25.730	26.104	25.897	25.645	25.938	25.833	25.873	25.288	23.051	23.668	23.567	23.303	23.040	23.508	23.161	23.088
Pre	Median	25	26	26	25	26	56	26	26	23	23	23	23	23	23	23	23
Pre	Brightness	504343	494926	511031	521085	530072	548040	540798	557372	461882	471499	464272	481975	452890	460961	471202	444522
	Pixels	19601	18960	19733	20319	20436	21214	20602	21698	20037	19921	19700	20683	19657	19609	20345	19253
Actual	Mass (ug)	0.035	0.07	0.105	0.14	29.323	58.506	87.689	116.872	0.035	0.07	0.105	0.14	29.323	58.506	87.689	116.872
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	Hand	22	ч	ĸ	æ	æ	æ	×	æ	ľ	ľ	L	ľ	ľ	L	Г	L
	Side	D	Ω	Ω	Ω	Ω	D	Ω	Ω	D	D	Ω	D	D	D	D	Ω

Table H.4. Simulated Exposures - Subject 17

Side Hand Actual Free Prea Preat Preat Preat Preat Preat Preat Preat Net Net Net Net Net Net Net Net Mass (ug) Pread Pread Pread Pread Pread Net Net Net Mass (ug) Pread Net Net<																					
Hand Actual Pee Pre Pre Post Post Post Ned Post Ned Ned <th< td=""><td>VITAE</td><td>Mass (ug)</td><td>0.088</td><td>0.081</td><td>0.089</td><td>1.475</td><td>2.243</td><td>3.066</td><td>7.085</td><td>12.939</td><td>19.917</td><td>0.228</td><td>0.110</td><td>0.119</td><td>0.351</td><td>1.104</td><td>1.192</td><td>2.872</td><td>4.243</td><td>7.849</td><td>11.774</td></th<>	VITAE	Mass (ug)	0.088	0.081	0.089	1.475	2.243	3.066	7.085	12.939	19.917	0.228	0.110	0.119	0.351	1.104	1.192	2.872	4.243	7.849	11.774
Hand Actual Pre Pre Pre Post Post Pref Net Hand Actual Mass (ug) Prode Brightness Median Avenage Brightness Prode <	Area	cm^2	116.11	119.02	121.77	123.42	117.54	118.25	123.49	118.48	118.04	115.34	112.06	110.06	107.13	109.57	109.59	106.59	110.88	109.43	105.58
Hand Actual Pre Pre Pre Prot	Net	Median	55	55	68	75	81	84	88	106	115	59	56	47	63	73	76	80	98	93	110
Hand Actual Pre Pre Pre Pre Pre Pre Pret Pret Prost	Net	Brightness	58388	55469	158190	286274	362218	458272	561549	680841	921040	130572	70419	82912	178782	316265	460016	\$39317	620298	792960	971541
Hand Actual Pre Pre Pre Post Post Post H and Mass (ug) Pixels Brightness Median Average Brightness Median R 1 0.052 18879 944850 53 50.048 957335 53 R 2 0.243 19353 967274 53 49.981 103903 54 R 4 2.712 19800 968674 52 48.923 103903 54 R 4 2.712 19800 96674 52 48.923 103903 54 R 5 17.693 20069 991433 52 49.402 119739 54 R 6 28.36 19113 958004 53 49.860 117973 56 R 7 28.958 19227 956094 52 49.630 117973 58 R 8 27.639 19194 958223	Net	Pixels	1074	1034	2432	3761	4246	860\$	6022	5813	7175	2217	1294	1665	2910	4114	5623	6146	6495	7369	8120
Hand Actual Pre Pre Pre Prost Hand Mass (ug) Pixels Brightness Median Average Brightness R 1 0.052 18879 944850 53 50.048 957535 R 1 0.052 18879 944850 53 49.981 981256 R 2 0.243 19353 567274 53 49.981 981256 R 4 2.712 19800 968674 52 48.923 1039903 R 4 2.712 19800 968674 52 49.981 107598 R 5 17693 20069 991453 52 49.402 1107598 R 6 28.36 19113 958664 53 49.860 1179531 R 7 28.958 19227 958664 53 49.360 1179531 R 8 27.639 19194 958223 5	Post	Average	50.709	50.732	52.528	55.195	58.559	61.363	63.689	71.326	80.374	48.511	48.050	47.435	49.582	53.866	58.396	62.293	64.760	72.927	82.133
Hand Actual Pre Pre Pre Hand Mass (ug) Pixels Brightness Median Average R 1 0.052 18879 944850 53 50.048 R 2 0.243 19353 967274 53 49.981 R 4 2.712 19800 968674 52 48.923 R 4 2.712 19800 968674 52 49.921 R 4 2.712 19800 968674 52 48.923 R 4 2.712 19800 968674 52 48.923 R 5 17.693 20069 991453 52 49.402 R 6 28.36 1913 95864 53 49.860 R 7 28.958 1927 958094 52 49.360 R 8 27.639 18754 860366 49 45.052 L <	Post	Median	53	53	54	54	56	57	58	58	9	51	50	49	51	51	54	56	55	57	59
Hand Actual Pre Pre Hand Mass (ug) Pixels Brightness Median R 1 0.052 18879 944850 53 R 2 0.243 19353 967274 53 R 4 2.712 19800 968674 53 R 5 17.693 20069 991453 52 R 6 28.36 1913 958707 53 R 7 28.958 19257 956094 52 R 8 27.639 20080 990861 52 L 1 0.052 18754 860366 49 L	Post	Brightness	957535	981256	1039903	1107598	1119231	1179757	1278617	1373734	1542370	915891	875422	848843	854600	959561	1040786	1079411	1167564	1297813	1410799
Hand Actual Pre Hand Mass (ug) Pixels Brightness R 1 0.052 18879 944850 R 2 0.243 19353 967274 R 2 0.243 19353 967274 R 2 0.243 19353 967274 R 4 2.712 19800 968674 R 5 17.693 20069 991453 R 6 28.36 19113 958707 R 8 27.639 20080 990861 R 8 27.639 20080 990861 R 8 27.639 20080 990861 L 1 0.052 18754 860366 L 2 0.398 18221 838260 L 3 2.458 17420 796852 L 4 5.981 17420 796852 L 5 <	Pre	Average	50.048	49.981	48.923	49.402	50.160	49.860	49.346	49.630	49.923	45.876	46.005	46.027	46.221	46.515	46.366	46.644	45.385	45.978	46.597
Hand Actual Pixels R 1 0.052 18879 R 2 0.243 19353 R 2 0.243 19353 R 4 2.712 19800 R 4 2.712 19800 R 5 17.693 20069 R 5 17.633 20069 R 6 28.36 19113 R 7 28.958 19227 R 9 93.917 19265 R 10 157.319 19194 L 1 0.052 18754 L 2 0.398 17806 L 4 5.981 17420 L 5 42.678 17820 L 6 72.086 17820 L 7 94.205 1732 L 9 174.403 17793 L 9 174.403 17168<	Pre	Median	53	53	52	52	53	53	52	52	52	49	49	49	49	50	49	50	49	49	50
Hand Actual R 1 0.052 R 2 0.243 R 2 0.243 R 4 2.712 R 4 2.712 R 5 17.693 R 6 28.36 R 7 28.958 R 8 27.639 R 9 93.917 R 9 93.917 L 1 0.052 L 1 0.052 L 3 2.458 L 4 5.981 L 5 42.678 L 6 72.086 L 7 94.205 L 9 174.403 L 9 174.403 L 10 214.988	Pre	Brightness	944850	967274	968674	991453	958707	958664	990861	956094	958223	860366	838260	823708	796852	828764	826237	808432	818288	818091	186661
Hand R 1 R 2 R 4 R 5 R 6 R 7 R 8 R 9 R 10 L 1 L 2 L 2 L 6 L 6 L 6 L 6 L 6 L 7 L 9 L 10		Pixels	18879	19353	19800	50069	19113	19227	20080	19265	19194	18754	18221	17896	17420	17817	17820	17332	18030	17793	17168
Hand	Actual	Mass (ng)	0.052	0.243	2.712	17.693	28.36	28.958	27.639	93,917	157.319	0.052	0.398	2.458	5.981	42.678	72.086	94.205	108.063	174.403	214.988
			-	2	4	\$	9	7	00	6	10	-	2	3	4	S	9	7	90	6	10
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		-	Ы	Ы	d.	Ы	Ы	Ь	Д	Ы	Ы	Ы	Ы	Ы	Ы	Д	Ъ	Ы	G,	Ы	Ъ

Table H.4. Continued

			_			_		,									
VITAE	Mass (ug)	0.094	0.156	0.144	0.217	20.777	38.513	49.826	60.219	960'0	0.098	0.091	0.120	17.682	38.487	49.773	57.357
Area	cm^2	121.47	121.41	119.53	125.52	123.12	125.31	128.15	127.01	106.27	102.85	104.46	112.58	116.79	120.35	119.89	115.46
Net	Median	31	33	33	32	121	168	205	227	17	32	30	32	96	177	215	228
Net	Brightness	45597	78933	63101	102238	1039608	1746234	2098417	2436138	73026	57834	61236	62420	955758	1748538	2151258	2440833
Net	Pixels	1332	2133	1602	2833	9292	11150	11919	13169	3165	1733	21114	1784	7931	10924	12194	13178
Post	Average	23.807	24.879	24.670	25.461	966.996	97.146	111.706	127.260	21.252	22.360	21.956	23.425	63.916	99.904	119.368	137.491
Post	Median	22	23	22	23	30	46	09	92	20	21	21	22	29	48	81	149
Post	Brightness	469567	491677	479515	520650	1341050	1977208	2326395	2628151	367264	374035	372946	429405	1214349	1955220	2326125	2581126
Pre	Average	22.769	22.559	22.858	22.550	22.250	22.033	22.413	21.945	20.830	20.963	20.978	21.706	21.399	21.478	21.424	21.656
Pre	Median	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21	21
Pre	Brightness	449740	445332	444263	460236	445455	448925	467046	453198	359935	350572	356329	397322	406382	420301	417664	406568
	Pixels	19752	19742	19436	20410	20020	20375	20838	20652	17280	16723	16986	18305	18991	19569	19495	18774
Actual	Mass (ug)	0.036	0.072	0.108	0.144	29.268	58.392	87.516	116.64	0.036	0.072	0.108	0.144	29.268	58.392	87.516	116.64
-	\dashv	-	7	9	4	S	9	7	~		7	3	4	S	9	7	00
	Hand	ж	æ	2	æ	м	ĸ	æ	æ	r	T	L	L	r	Г	L	L
	Side	Q	Δ	Ω	Ω	۵	Ω	D	Ω	Ω	Ω	۵	Ω	Ω	Ω	Ω	D

Table H.5. Simulated Exposures - Subject 18

Side Hund Actual Pre Pre Pre Pre Pre Pres Pr																						
Hand Actual Pre Pre Pres Pres Post Post Pres Pres <th< td=""><td>VITAE</td><td>Mass (ug)</td><td>0.075</td><td>0.115</td><td>0.595</td><td>1.274</td><td>3.112</td><td>4.880</td><td>6.916</td><td>8.490</td><td>16.352</td><td>21.685</td><td>0.144</td><td>0.295</td><td>0.514</td><td>0.825</td><td>1.807</td><td>3.115</td><td>4.557</td><td>5.896</td><td>11.331</td><td>16.105</td></th<>	VITAE	Mass (ug)	0.075	0.115	0.595	1.274	3.112	4.880	6.916	8.490	16.352	21.685	0.144	0.295	0.514	0.825	1.807	3.115	4.557	5.896	11.331	16.105
Hand Actual Pre Pre Pre Pre Pre Pre Pres Pre	Area	cm^2	88.82	89.92	92.80	88.82	94.89	93.63	87.21	89.29	94.32	90.22	94.03	94.27	91.01	89.70	91.08	92.76	88.81	98.98	83.26	86.30
Hand Actual Pres Pre Pres Pres <t< td=""><td>Net</td><td>Median</td><td>42</td><td>44</td><td>89</td><td>9/</td><td>91</td><td>102</td><td>106</td><td>109</td><td>129</td><td>139</td><td>52</td><td>54</td><td>58</td><td>63</td><td>72</td><td>78</td><td>85</td><td>68</td><td>107</td><td>112</td></t<>	Net	Median	42	44	89	9/	91	102	106	109	129	139	52	54	58	63	72	78	85	68	107	112
Hand Actual Pre Pre Pre Prost	Net	Brightness	81645	84582	215979	314178	471310	177555	656081	749089	927778	1048014	91177	170717	246767	308831	432833	514952	590550	686945	846823	1000921
Hand Actual Pre Pre Pre Pre Pre Pres Prest	Net	Pixels	2104	1863	3228	4015	4871	5161	5753	6429	0999	7042	1774	3112	4158	4682	5603	5945	6231	8069	7055	7854
Hand Actual Pre Pre Pre Pre Pre Prost Post R 1 0.073 14443 655863 47 45.410 622377 R 1 0.073 14443 655863 47 45.410 622377 R 2 0.369 14621 660086 47 45.146 622377 R 3 2.021 15090 682576 47 45.146 622377 R 4 7.167 14443 645072 47 44.663 76367 R 4 7.167 14443 645072 47 44.617 93517 R 4 7.167 14481 63304 47 44.617 93517 R 6 28.593 1451 671398 47 44.575 100759 R 7 39.99 14181 631076 47 44.575 100769 R 9 86.041	Post	Average	43.122	44.811	49.517	53.054	60.667	66.202	73.056	76.812	86.084	95.384	42.097	44.269	46.625	49.535	55.194	59.602	64.934	71.175	83.250	90.700
Hand Actual Pre Pre Pre Hand Mass (ug) Pixels Brightness Median Average R 1 0.073 14443 653863 47 45.410 R 2 0.369 14621 660086 47 45.146 R 3 2.021 15090 682576 47 45.234 R 4 7.167 14443 645072 47 44.663 R 4 7.167 14443 645072 47 44.663 R 5 20.919 15429 688392 47 44.617 R 6 28.593 15225 672598 47 44.575 R 8 44.408 14519 653948 47 44.574 R 9 86.041 15336 683901 47 44.594 R 10 120.697 14520 614769 42 40.105 L	Post	Median	45	46	48	49	90	52	55	57	57	59	44	46	46	48	50	50	52	54	57	59
Hand Actual Pre Pre Hand Mass (ug) Pixels Brightness Median R 1 0.073 14443 653863 47 R 2 0.369 14621 660086 47 R 3 2.021 15090 682576 47 R 4 7.167 14443 645072 47 R 4 7.167 14443 645072 47 R 4 7.167 14443 648072 47 R 4 7.167 14443 648072 47 R 5 20.919 15429 688392 47 R 6 28.593 14181 633948 47 R 9 86.041 15336 683901 47 R 9 86.041 15339 614769 47 L 1 0.073 15226 672844 42 L <td>Post</td> <td>Brightness</td> <td>622377</td> <td>654779</td> <td>747355</td> <td>766367</td> <td>935917</td> <td>1007798</td> <td>1039636</td> <td>1115310</td> <td>1320364</td> <td>1399288</td> <td>647710</td> <td>677496</td> <td>690055</td> <td>721181</td> <td>817261</td> <td>896716</td> <td>937644</td> <td>1005562</td> <td>1127124</td> <td>1271975</td>	Post	Brightness	622377	654779	747355	766367	935917	1007798	1039636	1115310	1320364	1399288	647710	677496	690055	721181	817261	896716	937644	1005562	1127124	1271975
Hand Actual Pre Hand Mass (ug) Pixels Brightness R 1 0.073 14443 655863 R 2 0.369 14621 660086 R 2 0.369 14621 660086 R 3 2.021 15090 682576 R 4 7.167 14443 645072 R 4 7.167 14443 645072 R 5 20.919 15429 688392 R 6 28.593 15225 672598 R 7 39.99 14181 633041 R 8 44.408 14519 653948 R 9 86.041 15336 683901 L 1 0.073 15290 612844 L 2 0.355 15329 614769 L 3 1.867 14799 602626 L 4 <	Pre	Average	45.410	45.146	45.234	44.663	44.617	44.177	44.575	45.041	44.594	44.381	40.081	40.105	40.721	40.493	39.349	39.752	39.967	41.119	41.330	41.255
Hand Actual Pixels R 1 0.073 14443 R 2 0.369 14621 R 2 0.369 14621 R 3 2.021 15090 R 4 7.167 14443 R 5 20.919 15429 R 6 28.593 14519 R 7 39.99 14181 R 8 44.408 14519 R 9 86.041 15326 L 1 0.073 15229 L 1 0.073 15290 L 3 1.867 14799 L 4 5.88 14586 L 5 23.324 14810 L 5 36.65 15083 L 7 40.069 14441 L 7 40.069 14424 L 9 75.421 13538	Pre	Median	47	47	47	47	47	47	47	47	47	47	42	42	43	43	43	43	43	43	43	43
Hand Aass (ug) R 1 0.073 R 2 0.369 R 3 2.021 R 4 7.167 R 5 20.919 R 6 28.593 R 7 39.99 R 7 39.99 R 8 44.408 R 9 86.041 R 10 120.697 L 1 0.073 L 2 0.355 L 3 1.867 L 5 23.324 L 5 23.324 L 6 36.65 L 7 40.069 L 8 38.44 L 9 75.421 L 9 75.421	Pre	Brightness	655863	660086	682576	645072	688392	672598	632117	653948	683901	6\$1076	612844	614769	602626	590628	582760	599579	\$77166	581748	559523	578892
Hand RR 1 RR 2 RR 3 RR 4 RR 7 RR 9 RR 10 L 1 1 L 2 L 2 L 3 L 6 L 6 L 6 L 6 L 7 L 9 L 9		Pixels	14443	14621	15090	14443	15429	15225	14181	14519	15336	14670	15290	15329	14799	14586	14810	15083	14441	14124	13538	14032
Hand Hand	Actual	Mass (ug)	0.073	0.369	2.021	7.167	20.919	28.593	39.99	44.408	86.041	120.697	0.073	0.355	1.867	5.88	23.324	36.65	40.069	38.44	75.421	97.982
╶╶╎┥┨┾╏┧┼╏┪╎╏╏ ┼┼┼┼			-	2	3	4	S	9	7	90	6	10	-	2	3	4	S	9	7	90	6	10
Sig a a a a a a a a a a a a a a a a a a a		Hand	R	×	×	ĸ	~	æ	æ	×	2	×	T	L	L	L	L	L	L	L	L	1
		Side	Ы	Ы	Д	Ь	Ч	Ь	Ы	Ь	Ь	Ч	Ы	Д	Ы	Д	ď	Ы	Ы	Ъ	Δ,	Δ,

Table H.5. Continued

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VITAE	Mass (ug)	0.045	0.053	0.094	6.562	14.054	22.791	29.741	32.683	0.047	0.054	0.133	5.652	13.439	22.178	30.612	36.803
Area	cm^2	86.41	85.41	19.68	89.46	91.89	92.53	88.68	89.54	93.91	85.19	91.81	94.94	95.47	98.52	94.99	99.76
Net	Median	22	23	41	106	121	144	168	179	21	26	29	79	108	126	160	189
Net	Brightness	69822	62274	62093	553979	880564	1186153	1333523	1422263	46754	43611	94616	552469	929346	1229568	1432592	1616853
Net	Pixels	3066	2452	1739	5043	6856	8170	8276	8595	1960	1486	2889	6125	7947	9035	9047	6096
Post	Average	27.429	28.341	29.494	58.456	75.916	93.233	106.165	110.695	25.478	27.554	28.479	53.071	73.936	89.712	104.918	113.551
Post	Median	26	26	28	35	42	52	63	29	24	26	27	34	42	53	62	69
Post	Brightness	385300	393937	429377	850066	1131453	1405765	1525703	1612156	388674	381681	425166	819409	1148822	1436917	1626652	1803182
Pre	Average	29.321	29.246	28.914	29.555	29.072	29.114	29.347	29.400	25.626	26.659	25.431	25.420	24.938	24.876	25.424	25.450
Pre	Median	28	28	28	28	28	28	28	28	25	26	25	25	25	25	25	25
Pre	Brightness	412001	406169	421300	429936	434372	438021	423179	428070	391307	369282	379666	392441	387120	398486	392704	404140
	Pixels	14051	13888	14571	14547	14941	15045	14420	14560	15270	13852	14929	15438	15523	16019	15446	15880
Actual	Mass (ug)	0.034	0.068	0.102	14.912	29.722	44.53	75.845	107.158	0.034	0.068	0.102	14.912	29.722	44.53	75.845	107.158
		-	7	3	4	\$	9	7	∞	-	2	3	4	\$	9	7	90
	Hand	Я	R	R	22	æ	~	R	R	L	Г	r	Г	r	L	L	ľ
	Side	D	D	D	D	D	D	D	D	D	D	D	D	D	D	D	D